SEARCH REQUEST FORM

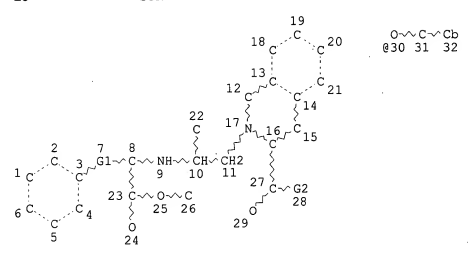
Scientific and Technical Information Center

Requester's Full Name: ZINNA N. DAVIS Examiner #: 65429 Date: 4/17/ccc2 Art Unit: 1625 Phone Number 30 8 4679 Serial Number: 09/424,673 Mail Box and Bldg/Room Location: (N13.3A07 Results Format Preferred (circle): PAPER DISK E-MAIL
If more than one search is submitted, please prioritize searches in order of need.
Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched. Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or utility of the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc, if known. Please attach a copy of the cover sheet, pertinent claims, and abstract.
Title of Invention: 2 Monsalvatje et &
Title of Invention: Monsalvate et al Inventors (please provide full names): Process for propuring quinquight
Earliest Priority Filing Date: 5/29/1997
For Sequence Searches Only Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the appropriate serial number.
Process for Preparing 7 10 10 10 10 10 10 10 10 10
Point of Contact: Beverly Shears Technical Info. Specialist CM1 1E05 Tel: 308-4994

STAFF USE ONLY Searcher: Bowerly 64964	Type of Search NA Sequence (#)	Vendors and cost where applicable
Searcher Phone #:	AA Sequence (#)	Dialog
Searcher Location:	Structure (#)	Questel/Orbit
Date Searcher Picked Up:	Bibliographic	Dr.Link
Date Completed: 04-21-03	Litigation	Lexis/Nexis
Searcher Prep & Review Time: 12	Fulltext	Sequence Systems
-al Prep Time:	Patent Family	WWW/Internet
ime:20	Other .	Other (specify)
30 (0.00)		

09/424673

(FILE 'REGISTRY' ENTERED AT 11:51:05 ON 21 APR 2003) L5 STR



REP G1=(1-3) CH2 VAR G2=OH/30 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 32

100.0% PROCESSED 4015 ITERATIONS SEARCH TIME: 00.00.01

0 ANSWERS

rs

STR

REP G1=(1-3) CH2 VAR G2=OH/30 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM GGCAT IS UNS AT 32 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 32

STEREO ATTRIBUTES: NONE

L10 67 SEA FILE=REGISTRY SSS FUL L8

100.0% PROCESSED 421 ITERATIONS

SEARCH TIME: 00.00.01

FILE 'HCAPLUS' ENTERED AT 12:01:30 ON 21 APR 2003

L11 25 S L10/P

L12 15 S L11 NOT (PY=>1997 OR PD=>19970529)

E1 THROUGH E38 ASSIGNED

L12 ANSWER 1 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1995:388940 HCAPLUS

DOCUMENT NUMBER:

123:111822

TITLE:

ACE inhibitors as a template for the design of

67 ANSWERS

bradykinin B2 receptor antagonists

AUTHOR(S):

Hoyer, Denton; Awad, Mohamed M. A.; Salvino, Joseph M.; Seoane, Peter R.; Dolle, Roland E.;

Houck, Wayne T.; Sawutz, David G.

CORPORATE SOURCE:

Department of Medicinal Chemistry, Sterling Winthrop Pharmaceutical Research Division,

Collegeville, PA, 19426-0900, USA

SOURCE:

Bioorganic & Medicinal Chemistry Letters (1995),

5(4), 367-70

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER:

Elsevier

DOCUMENT TYPE: Journal LANGUAGE: English

Angiotensin converting enzyme (ACE) degrades both angiotensin II and AB bradykinin. Accordingly, it was hypothesized that ACE inhibitors can serve as models to design antagonists for the bradykinin receptor. The potent ACE inhibitor quinapril was modified to serve as a spacer sepg. two lipophilic pos. charges required for bradykinin binding. The synthesis and bradykinin receptor-binding

activity of a series of antagonists (quinapril derivs.) based on

this hypothesis were described.

ΙT 85441-61-8DP, Quinapril, analogs RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(prepn. of quinapril derivs. as bradykinin B2 receptor antagonists)

85441-61-8 HCAPLUS RN

3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-CN phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L12 ANSWER 2 OF 15 HCAPLUS COPYRIGHT 2003 ACS

1992:572104 HCAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 117:172104

TITLE: Methods for the synthesis of aminosuberic acid

derivatives

Hoffmann, Gerhard; Liedtke, Bernhard; Vollmer, INVENTOR(S):

Karl Otto

PATENT ASSIGNEE(S): Goedecke AG, Germany SOURCE: Ger. Offen., 6 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC: NUM. COUNT:

PATENT INFORMATION:

PATENT NO. APPLICATION NO. DATE KIND DATE A1 19920604 DE 1990-4037960 19901129 DE 4037960 PRIORITY APPLN. INFO.: DE 1990-4037960 19901129

CASREACT 117:172104; MARPAT 117:172104 OTHER SOURCE(S):

For diagram(s), see printed CA Issue. GT

AΒ Aminosuberic acid derivs. I (R1 = alkyl or alkenyl with up to 6

> 308-4994 Searcher : Shears

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carbon atoms, C5-7-cycloalkyl, C5-7-cycloalkenyl, C7-12-cycloalkylalkyl, C6-10-aryl, C7-14-aralkyl, mono- or bicyclic heterocyclic group; R2 = aryl; R3 = C1-6-alkyl, C2-6-alkenyl, C7-14-aralkyl; Z forms a heterocyclic ring) were prepd. by the selective cleavage of PhCH2O2CCHR1NHCH(CO2R3)CH2COR2 with AlC13, condensing the resulting HO2CCHR1NHCH(CO2R3)CH2COR2 with heterocyclic compd. II, and hydrogenating the resulting ketone III with H2, deuterium or tritium. Thus, (S,S)-PhCOCH2CH(CO2Et)-Ala-OCH2Ph was debenzylated with AlC13 in the presence of anisole in CH2C12/MeNO2 to give 86% (S,S)-PhCOCH2CH(CO2Et)-Ala-OH, which was condensed with (S)-1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid benzyl ester by DCC/1-hydroxybenzotriazole in CH2C12 to give product IV (isolated as the HCl salt). IV was hydrogenated over Pd/C in the presence of HCl to give 88% tetrahydroisoquinoline V.HCl.

IT 82586-55-8P 143381-51-5P

RN 82586-55-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 143381-51-5 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl-3,3-d2]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, [3S-[2[R*(R*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

HCAPLUS COPYRIGHT 2003 ACS L12 ANSWER 3 OF 15

ACCESSION NUMBER: 1991:207829 HCAPLUS

DOCUMENT NUMBER: 114:207829

Preparation of carboxyalkyl dipeptides useful as TITLE:

angiotensin-converting enzyme (ACE) inhibitors

INVENTOR(S): Oudenes, Jan; Schleicher, Richard Henry

PATENT ASSIGNEE(S): Pharma Investi S. A., Spain

SOURCE:

Span., 10 pp. CODEN: SPXXAD

DOCUMENT TYPE: Patent

LANGUAGE: Spanish

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ES 2004804	A6	19890201	ES 1987-2390	19870813
PRIORITY APPLN.	INFO.:	ES	1987-2390	19870813 ⁻
ARIJER AAJJRAE (A)	1.47	DD3M 114 007000		

OTHER SOURCE(S): MARPAT 114:207829

 R^1

GI

R1R2R3CNHCHRCONR4CHR5COR6 [R, R1, R2 = H, alkyl, Ph, phenylalkyl, alkylphenyl, aminoalkyl, protected aminoalkyl; R3 = CO2H or its ester; R4 = H, alkyl; R5 = H, alkyl, Ph, phenylalkyl, alkylphenyl; R4R5 may form (un) substituted C4-9 monocyclic or fused bicyclic nucleus; R6 = OH, alkoxy, alkenyloxy, OPh, alkylsilyloxy, etc.], including such ACE inhibitors as enalapril, lisinopril, indolapril, ramipril, and quinapril, were prepd. by converting carboxylakyl R1R2R3CNHCHRCO2H to cyclic anhydrides I, reaction of I with

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R4NHCHR5COR6, and optional deprotection, sapon. of R6, or salification. Thus, N-(1-S-ethoxycarbonyl-3-phenylpropyl)-L-alanine was treated with 1,1-carbonyldiimidazole in EtOAc at 20.degree., followed by L-proline. Two crystns. with maleic acid gave first imidazole maleate byproduct and then 77% 1-[N-[(S)-1-(ethoxycarbonyl)-3-phenylpropyl]-L-alanyl]-L-proline, i.e. enalapril maleate.

IT **85441-61-8P**, Quinapril

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, via cyclic anhydride)

RN 85441-61-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L12 ANSWER 4 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1989:38911 HCAPLUS

DOCUMENT NUMBER: 110:38911

TITLE: Preparation of crystalline quinapril, a known

antihypertensive

INVENTOR(S): Goel, Om P.; Krolls, Uldis PATENT ASSIGNEE(S): Warner-Lambert Co., USA

SOURCE: U.S., 4 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

			•	
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4761479	A	19880802	US 1987-32209	19870330
AU 8812383	A1	19880929	AU 1988-12383	19880229
AU 605555	B2	19910117		
ZA 8801426	Α	19891025	ZA 1988-1426	19880229
CA 1291999	A1	19911112	CA 1988-560594	19880304
EP 285992	A1	19881012	EP 1988-105131	19880329
EP 285992	B1	19910403		,
R: AT, BE,	CH, DE	, ES, FR,	GB, GR, IT, LI, LU, NL	, SE
JP 63258459	A2	19881025	JP 1988-73492	19880329
AT 62229	E	19910415	AT 1988-105131	19880329
PRIORITY APPLN. INFO	. :		US 1987-32209 EP 1988-105131	19870330 19880329
			EP 1900-103131	19000329

09/424673

AB The title compd. (I) an angiotensin converting enzyme and antihypertensive agent is prepd. in a highly pure state. A soln. of I in glacial AcOH contg. HCl(g) was stirred for 2 h 20 min and was dild. with xylene; the process was repeated, the glossy solid dissolved in MeNC at 60.degree., and the product dried at 50.degree. under vacuum for 16 h to give 91.2% I-HCl.

TT 82586-55-8P, Quinapril hydrochloride RL: SPN (Synthetic preparation); PREP (Preparation)

(cryst., prepn. of) RN 82586-55-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

L12 ANSWER 5 OF 15 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1986:608744 HCAPLUS

ACCESSION NUMBER:
DOCUMENT NUMBER:

105:208744

TITLE:

Synthesis of novel angiotensin converting enzyme inhibitor quinapril and related compounds. A

divergence of structure-activity relationships

for non-sulfhydryl and sulfhydryl types

AUTHOR(S):

Klutchko, Sylvester; Blankley, C. John; Fleming, Robert W.; Hinkley, Jack M.; Werner, Ann E.; Nordin, Ivan; Holmes, Ann; Hoefle, Milton L.;

Cohen, David M.; et al.

CORPORATE SOURCE:

Dep. Chem., Warner-Lambert/Parke-Davis Pharm.

Res., Ann Arbor, MI, 48106, USA

SOURCE:

Journal of Medicinal Chemistry (1986), 29(10),

1953-61

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 105:208744

GI

Ι

AΒ The synthesis and angiotensin-converting enzyme (ACE) inhibiting activities of quinapril (S,S,S)-I (R=R2=H,R1=Et), its active diacid (S,S,S)-I (R=R1=R2=H), and its dimethoxy analog (S,S,S)-I (R = H, R1 = Et, R2 = MeO) are reported. Thus, (S)-1,2,3,4-tetrahydro-3-isoquinolinecarboxylic acid 1,1-dimethylethyl ester was acylated with (S,S)-PhCH2CH2CH(CO2Et)NHCHMeCO2H followed by hydrolysis of the product to give (S,S,S)-I (R=R2=H,R1=Et). These tetrahydro-3isoquinolinecarboxylic acid derivs. possess in vitro potency and in vivo efficacy equiv. to that of enalapril. Sulfhydryl analogs with the same structural variation are also highly potent. In contrast, tetrahydro-1-isoquinolinecarboxylic acid and homologous isoindoline-1-carboxylic acid analogs show a striking divergence in potency between the two types, sulfhydryl analogs being essentially inactive, while non-sulfhydryl analogs are equipotent with the proline prototype. This is the first evidence suggesting that alternate binding modes may exist for the two major structural classes of small mol. ACE inhibitors.

IT 82586-52-5P 82637-57-8P 85441-61-8P 89300-89-0P 103775-06-0P 103775-09-3P 103775-10-6P 103775-11-7P 103775-12-8P 103833-14-3P 103833-15-4P 103833-16-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and angiotensin converting enzyme inhibition activity of)

RN 82586-52-5 HCAPLUS

CN

3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 82637-57-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, phenylmethyl ester, [3S-[2[R*(R*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 85441-61-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 89300-89-0 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, [3S-[2[R*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

#C1

RN 103775-06-0 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, [3R-[2[S*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

HCl

RN 103775-09-3 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, [3S-[2[R*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 103775-10-6 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 103775-11-7 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, [3S-[2[R*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 103775-12-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, [3R-[2[S*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 103833-14-3 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, monohydrochloride, [3S-[2[R*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 103833-15-4 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, monohydrochloride, [3R-[2[S*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

HCl

RN 103833-16-5 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, [3R-[2[S*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 82586-55-8P 82637-58-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and angiotensin-converting enzyme inhibition activity of) 82586-55-8 HCAPLUS

RN 82586-55-8 HCAPLUS
CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-,
monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 82637-58-9 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, phenylmethyl ester, (3S)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 82637-57-8 CMF C34 H40 N2 O7

Absolute stereochemistry.

CRN 110-16-7 CMF C4 H4 O4

Double bond geometry as shown.

IT 82586-54-7P 103775-05-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and debenzylation of)

RN 82586-54-7 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, (3S)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 82586-53-6 CMF C32 H36 N2 O5

Absolute stereochemistry. Rotation (-).

CRN 110-16-7 CMF C4 H4 O4

Double bond geometry as shown.

RN 103775-05-9 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1R)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, (3S)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 103775-04-8 CMF C32 H36 N2 O5

Absolute stereochemistry.

CRN 110-16-7 CMF C4 H4 O4

Double bond geometry as shown.

L12 ANSWER 6 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1985:560859 HCAPLUS

DOCUMENT NUMBER:

103:160859

TITLE:

N-Alkylated dipeptides and their esters

INVENTOR(S):

Teetz, Volker; Wissmann, Hans; Urbach, Hansjoerg

PATENT ASSIGNEE(S):

Hoechst A.-G. , Fed. Rep. Ger.

SOURCE:

Eur. Pat. Appl., 33 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent German

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT NO.		KIND	DATE		APPLICATION NO.	DATE
ΕP	135182		A2	19850327		EP 1984-110678	19840907
ΕP	135182		A3	19860305			
ΕP	135182		B1	19880727			
	R: AT,	BE,	CH, DE	, FR, GB,	IT,	LI, LU, NL, SE	
DE	3333454		A1	19850411		DE 1983-3333454	19830916
ΑT	35997		Ε	19880815		AT 1984-110678	19840907
HU	36145		0	19850828		HU 1984-3415	19840910
ΗU	201565		В	19901128			
FI	8403590		Α	19850317		FI 1984-3590	19840913
FI	80464		В	19900228			

FI	80464	С	19900611			
CA	1338163	A1	19960312	C	A 1984-463078	19840913
DK	8404405	Α	19850317	D:	K 1984-4405	19840914
DK	164939	В	19920914			
DK	164939	С	19930201			4
NO	8403662	Α	19850318	N	0 1984-3662	19840914
NO	167743	В	19910826			
NO	167743	С	19911204			
AU	8433070	A1	19850321.	A	U 1984-33070	19840914
AU	576782	B2	19880908			
JP	60089497	A2	19850520	J.	P 1984-191868	19840914
JP	07098835	B4	19951025			
ZA	8407257	A	19850529	Z	A 1984-7257	19840914
ES	535917	A1	19851001	E	S 1984-535917	19840914
IL	72947	A1	19890228	I.	L 1984-72947	19840914
US	5068351	Α	19911126	U	S 1990-560004	19900727
PRIORITY	Y APPLN. INFO.:			DE 1	983-3333454	19830916
				EP 1	984-110678	19840907
				US 1	984-650715	19840914
				US 1	986-943882	19861219
				US 1	988-178767	19880330
				US 1	989-403919	19890907

GI

AΒ Title compds. R3O2CCHR4NR5COCHR1NHCH(CO2R2)(CH2)nR [n = 1, 2; R = H,](un) substituted C1-8 aliph., C3-9 alicyclic, C6-12 arom., C7-14 araliph., or C7-14 alicyclic aliph. residue, OR6, SR6 [R6 = (un)substituted C1-4 aliph., C6-12 arom., or heteroarom. residue]; R1 = H, (un)usbsituted C1-6 aliph., C3-9 alicyclic, C4-13 alicyclic aliph., C6-12 arom., C7-16 araliph., or heteroarom. residue, amino acid side chain; R2, R3 = H, (un)substituted C1-6 aliph., C3-9 alicyclic, C6-12 arom., or C7-16 araliph. residue; CHR4NR5 = C5-15 heterocyclic mono-, bi-, or tricyclic ring system] were prepd. via the condensation of HO2CCHR1NHCH(CO2R2)(CH2)nR with R3O2CCHR4NHR5 in the presence of phosphinic acid anhydrides R7R8P(0)OP(0)R9R10 (R7, R8, R9, R10 = alkyl or aralkyl). Thus, (S,S,S)azabicyclo[3.3.0]octane II was condensed with (S)-PhCH2CH2CH(CO2Et)-(S)-Ala-OH by ethylmethylphosphinic acid anhydride in CH2Cl2 contg. Et3N to give peptide III (R11 = CH2Rh), which was debenzylated to give III (R1 = H). I inhibit angiotensin-converting enzyme and can be used as antihypertensives (no data).

IT 82586-53-6P

RN 82586-53-6 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L12 ANSWER 7 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1985:560858 HCAPLUS

DOCUMENT NUMBER:

103:160858

TITLE:

N-Alkylated dipeptides and their esters

INVENTOR(S):

Urbach, Hansjoerg; Henning, Rainer; Wissmann,

Hans; Teetz, Volker

PATENT ASSIGNEE(S):

Hoechst A.-G. , Fed. Rep. Ger.

SOURCE:

Eur. Pat. Appl., 32 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	TENT NO.		KIND	DATE		API	PLICATION NO.	DATE
ΕP	135181 135181 135181		A2 A3 B1	19850327 19860402 19900131		EP	1984-110677	19840907
	R: AT,	ΒE,	CH, DE	, FR, GB,	IT,	LI, I	LU, NL, SE	
DE	3333455		A1				1983-3333455	
AT	49979		E				1984-110677	
HU	36140		0			HU	1984-3417	19840910
HU	198303		В	19890928				
ΓI	8403591		A			FI	1984-3591	19840913
FΙ	80275	٠	В	19900131				
FI	80275		С	19900510				
CA	1338162		· A1	19960312		CA	1984-463071	and the second s
DK	8404404		Α	19850317		DK	1984-4404	19840914
DK	166027		В	19930301				
DK	166027		С	19930712				
NO	8403663		A	19850318		NO	1984-3663	19840914
NO	167808		В	19910902				
NO	167808		C	19911218				
ΑU	8433071		A1	19850321		AU	1984-33071	19840914
ΑU	575585		B2	19880804			•	•
JΡ	60089498		A2	19850520		JP	1984-191869	19840914
JΡ	07098836		B4	19951025				

ZA 8407259		A	19850529		ZA 1984-7259	19840914
ES 535918		A1	19851001		ES 1984-535918	19840914
IL 72946		A1	19900429		IL 1984-72946	19840914
US 5055591		A	19911008		US 1988-173024	19880323
PRIORITY APPLN.	INFO.:			DE	1983-3333455	19830916
				EP	1984-110677	19840907
	•			US	1984-650714	19840914
				US	1986-943881	19861219

GΙ

Title compds. R3O2CCHR4NR5COCHR1NHCH(CO2R2)(CH2)nR [I; n = 1, 2; R = 1, 2;AB H, (un) substituted C1-8 aliph., C3-9 alicyclic, C6-12 arom., C7-14 araliph., or C7-14 alicyclic aliph. residue, OR6, SR6 [R6 = (un) substituted C1-4 aliph., C6-12 arom., or heteroarom. residue]; R1 = H, (un)substituted C3-9 alicyclic, C4-13 alicyclic aliph., C6-12 arom., C7-16 araliph., or heteroarom. residue, amino acid side chain; R2, R3 = H, (un) substituted C1-6 aliph., C3-9 alicyclic, C6-12 arom., or C7-16 araliph. residue; CHR4NR5 = C5-15 heterocyclic mono-, bi-, or tricyclic ring system] were prepd. via the condensation of HO2CCHR1NHCH(CO2R2)(CH2)nR with R3O2CCHR4NHR5 in the presence of an alkanephosphoric acid anhydride. Thus, (S,S,S)-azabicyclo[3.3.0]octane II was condensed with (S)-PhCH2CH2CH(CO2Et)-(S)-Ala-OH by n-propanephosphonic acid anhydride in CH2C12 in the presence of N-ethylmorpholine to give peptide deriv. III (R7 = CH2Ph), which was debenzylated to give III (R7 = H) (all-S isomer). I inhibit angiotensin-converting enzyme and can be used as antihypertensives (no data).

IT 82586-53-6P

RN 82586-53-6 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L12 ANSWER 8 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1984:175294 HCAPLUS

DOCUMENT NUMBER: 100:175294

Carboxyalkyl dipeptides and pharmaceutical TITLE:

compositions containing them

Smith, Elizabeth M.; Witkowski, Joseph T.; Doll, INVENTOR(S):

Ronald J.; Gold, Elijah H.; Neustadt, Bernard

R.; Yehaskel, Albert S.

PATENT ASSIGNEE(S):

Schering Corp., USA Eur. Pat. Appl., 134 pp. SOURCE:

CODEN: EPXXDW Patent

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 88350 EP 88350	A1 B1	19830914 19850220	EP 1983-102014	19830302 .
R: AT, BE,	CH, DE	, FR, IT, LI,	LU, NL, SE	
US 4431645	A	19840214	US 1982-355639	19820308
		19840214		19820308
ZA 8300362		19840926		19830119
AT 11921	E	19850315	AT 1983-102014	19830302
NO 8300737	Α	19830909	NO 1983-737	19830303
AU 8312035	A1	19830915	AU 1983-12035	19830303
AU 557795	B2	19870108		
GB 2117777	A1	19831019	GB 1983-5837	19830303
GB 2117777	B2	19850626		
ES 520261	A1	19840401	ES 1983-520261	19830303
DK 8301101	Α	19830909	DK 1983-1101	19830304
JP 58162561	A2	19830927	JP 1983-35707	19830304
FI 8300752	Α	19830909	FI 1983-752	19830307
ни 29605	0	19840228	ни 1983-781	19830307
HU 195520	В	19880530		
ZA 8301844	Α	19840627	ZA 1983-1844	19830316
PRIORITY APPLN. INFO	. :		US 1982-355638	19820308
			US 1982-355639	19820308

US 1982-360532 19820322 ZA 1983-362 19830119 EP 1983-102014 19830302

CASREACT 100:175294 OTHER SOURCE(S):

For diagram(s), see printed CA Issue. GI

Title compds. RCH2CR1(CO2H)-NHCH[(CH2)nXR2]CO-X1-OH[R = alkyl,AΒ PhCH2, PhCH2O, PhCH2S, PhO, PhS; R1 = H, alkyl; X = S, R2 = substituted (3,4-dihydro-7-sulfamoyl-1,2,4-benzothiadiazin-3-yl 1,1-dioxide) methyl; X = NR3 (R3 = H, alkyl, Ph), R2 = sulfamoyl-substituted Bz, PhSO2, or benzyl; XR2 = sulfamoyl-substituted N-contg. heterocyclic ring; n = 1-6; X1 =(un) substituted Pro or related N-contg. heterocyclic amino acid residues] were prepd. as antihypertensives and agents for the treatment of congestive heart failure and glaucoma (no data). Thus. H-L-Lys(Z)-OH (Z = CO2CH2Ph) was treated with PhCH2CH2COCO2Et and NaBH3CN to give (S)-PhCH2CH2CH(CO2Et)-L-Lys(Z)-OH, which was condensed with indole I to give dipeptide II (R4 = Z, R5 = CH2Ph), which was deblocked by hydrogenolysis to give II (R4 = R5 = H), which was sulfonylated with 4-chloro-3-sulfamoylbenzenesulfonyl chloride to give title compd. III.

IT 89083-69-2P

> RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

89083-69-2 HCAPLUS RN

CN 3-Isoquinolinecarboxylic acid, 7-[[3-(aminosulfonyl)-4chlorobenzoyl]amino]-2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]- $1-\exp(-1, 2, 3, 4-\text{tetrahydro-}, [3S-[2[R*(R*)], 3R*]]-(9CI)$ INDEX NAME)

Absolute stereochemistry.

HCAPLUS COPYRIGHT 2003 ACS L12 ANSWER 9 OF 15

ACCESSION NUMBER:

1984:138974 HCAPLUS

DOCUMENT NUMBER:

100:138974

TITLE:

SOURCE:

Tetrahydroisoquinoline derivatives

PATENT ASSIGNEE(S):

Takeda Chemical Industries, Ltd., Japan

Jpn. Kokai Tokkyo Koho, 8 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

Searcher

Shears

308-4994

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 58188857 A2 19831104 JP 1982-69898 19820426
PRIORITY APPLN. INFO.: JP 1982-69898 19820426
GI

AB Tetrahydroisoquinoline derivs. I [R, R1, configuration = Me3C, Me3C, R (II); Me3C, Me3C, S (III); Me3C, H, R (HBr); Me3C, H, S (HBr); H, Me3C, R; H, Me3C, S; H, H, R (HCl); H, H, S (HCl)] were prepd. by reaction of IV with PhCH2CH2COCO2Et (V) optionally followed by hydrolysis. I had angiotensin-converting enzyme inhibitory and bradykinin-decompg. enzyme inhibitory activities and are useful as hypotensives (no data). Thus, a mixt. of (3S)-IV oxalate (R = Me3C) 3.2, AcOH 1.6, NaOAc 0.6, V 3.5, Mol. Sieve 3A 6, and Raney Ni 5 g in EtOH was hydrogenated to give 0.5 g II and 0.7 g III.

IT 82586-55-8P 89245-28-3P 89300-87-8P 89300-89-0P

RN 82586-55-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 89245-28-3 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 7-(1,1-dimethylethyl)-2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrobromide, [3S-[2[R*(S*)],3R*]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

• HBr

RN 89300-87-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 7-(1,1-dimethylethyl)-2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrobromide, [3S-[2[R*(R*)],3R*]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

• HBr

RN 89300-89-0 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, [3S-[2[R*(S*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

HC1

HCAPLUS COPYRIGHT 2003 ACS L12 ANSWER 10 OF 15

1983:216005 HCAPLUS ACCESSION NUMBER:

98:216005 DOCUMENT NUMBER:

Tetrahydroisoquinolines TITLE:

PATENT ASSIGNEE(S): Tanabe Seiyaku Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 13 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

			
JP 58021666 A2 19	9830208	JP 1981-120547	19810730
PRIORITY APPLN. INFO.:	JP	1981-120547	19810730

AΒ I (R = alkyl; R1 = alkyl, phenylalkyl; R2, R3 = H, alkyl,phenylalkyl) were prepd. by acylation of II (R4 = alkyl, phenylalkyl; R5 = H) with HO2CCHRX (X = halo) and condensation of the resulting II (R5 = COCHRX) with H2NCHR1CO2R6 (R6 = alkyl, phenylalkyl) followed by deesterification and optional esterification. Thus, a mixt. of 8.0 g II (R4 = CH2Ph, R5 = COCHBrMe), prepd. from HO2CCHMeBr and II (R4 = CH2Ph, R5 = H), 20 mL P(NMe2)3, 2.8 g K2CO3, and 5.1 g L-H-Phe-OCH2Ph was stirred at room temp. for 1 day to give 4.8~g .alpha.- and 1.0~g.beta.-stereoisomers of I (R = Me, R1 = R2 = R3 = CH2Ph). Hydrogenation of 2.0 g the .alpha.-stereoisomer over Pd/C gave 1.0 g I (R2 = R3 = H). I inhibited angiotensin-converting enzyme in vitro.

IT 85892-46-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as antihypertensive agent)

RN 85892-46-2 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 1,2,3,4-tetrahydro-2-[1-oxo-2-[[2-oxo-2-(phenylmethoxy)-1-(phenylmethyl)ethyl]amino]propyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

L12 ANSWER 11 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1983:179903 HCAPLUS

DOCUMENT NUMBER:

98:179903

TITLE:

Isoquinolinecarboxylic acid derivatives and

pharmaceutical composition containing them

INVENTOR(S):

Patchett, Arthur A.; Wu, Mu Tsu

PATENT ASSIGNEE(S):

Merck and Co., Inc. , USA

SOURCE:

Eur. Pat. Appl., 30 pp. CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 65301	A1	19821124	EP 1982-104291	19820517
R: AT, BE,	CH, DE	, FR, GB,	IT, LU, NL, SE	
JP 58004770	A2	19830111	JP 1982-82521	19820518
PRIORITY APPLN. INFO.	:	•	US 1981-264880	19810518
			US 1982-362082	19820329
GI				

AB Title compds. I [R, R4 = H, alkyl, aralkyl; R1 = C1-10 alkyl, substituted C1-6 alkyl, (un)substituted aralkyl or heteroaralkyl; R2 = H, (un)substituted C1-6 alkyl; R3 = H, halo, alkoxy] were prepd. as angiotensin-converting enzyme inhibitors and antihypertensives (no data). Thus, Z-Ala-OH (Z = PhCH2O2C) was condensed with isoquinoline II by DCC to give a product, which was sapond. and then Z-deblocked by HBr/HOAc to give alanylisoquinoline III (R5 = H), which was condensed with PhCH2CH2COCO2Et in the presence of NaBH3CN to III [R5 = CH(CO2Et)CH2CH2Ph].

IT 82768-84-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and sapon. of)

RN 82768-84-1 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro- (9CI) (CA INDEX NAME)

IT 85441-61-8P

RN 85441-61-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L12 ANSWER 12 OF 15 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1983:89930 HCAPLUS

09/424673

DOCUMENT NUMBER:

98:89930

TITLE:

Amidoamino acids and pharmaceutical preparations

containing them

INVENTOR(S):

Suh, John T.; Barton, Jeffrey N.; Regan, John R.

USV Pharmaceutical Corp., USA

SOURCE:

Belg., 27 pp. CODEN: BEXXAL

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT ASSIGNEE(S):

P	ATENT NO.	KIND	DATE	I	APPLICATION NO.	DATE
	E 892552 N 156096	A1 A	19820920 19850511		BE 1982-207610 IN 1982-CA265	19820318 19820308
_	L 8201066	A	19821018		NL 1982-1066	19820315
-	L 65247	A1	19870731		IL 1982-65247	19820315
S	E 8201654	Α	19820920		SE 1982-1654	19820316
A	U 8281584	A1	19820923	7	AU 1982-81584	19820316
A	U 558451	B2	19870129			
G	B 2095252	Α	19820929	. (GB 1982-7770	19820317
G	B 2095252	B2	19850417			
D	E 3209708	· A1	19821021		DE 1982-3209708	19820317
E	S 510498	A1	19830801		ES 1982-510498	19820317
N	0 8200903	Α	19820920		NO 1982-903	19820318
D	K 8201209	Α	19820920	_	DK 1982-1209	19820318
	P 57165355	A2	19821012		JP 1982-41801	19820318
Z	A 8201833	Α	19830126	_	ZA 1982-1833	19820318
C	Н 658455	Α	19861114		CH 1982-1691	19820318
F	I 8200974	Α	19820920	_	FI 1982-974	19820319
_	R 2502149	A1	19820924]	FR 1982-4738	19820319
	R 2502149	В1	19850118			
	S 521030	A1	19840516	_	ES 1983-521030	19830316
	S 521031	A1	19841201		ES 1983-521031	19830316
	TY APPLN. INFO.	:		US :	1981-245407	19810319
GI						

$$R^{7}O_{2}C$$
 $(CH_{2})_{m}$
 $R^{8}n$
 $R^{1}O_{2}CCR^{2}R^{3}NR^{4}CR^{5}R^{6}CON$
 $(CH_{2})_{m}$
 I
 $HO_{2}C$
 $PhCH_{2}CH_{2}CH(CO_{2}H)-Ala-N$

Amino acid-substituted tetrahydroisoquinoline derivs. and related compds. I [R1, R7 = H, alkyl, phenylalkyl; R2-R6 = H or ΑB (un) substituted alkyl, alkenyl, alkynyl, aryl, cycloalkyl, or

Searcher :

Shears

308-4994

09/424673

heterocyclyl; R8 = alkyl, NO2, amino, OH, halo, CF3, etc.; m = 0-2, m' = 1-3, n = 0-4] and their salts were prepd. The products are useful as antihypertensives (no data). Thus, Me L-1,2,3,4-tetrahydro-3-isoquinolinecarboxylate HCl salt was acetylated with PhCH2O2C-Ala-OH and the product deprotected and then treated with PhCH2CH2COCO2H followed by redn. with NaBH3CN to give II.

IT 82586-53-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and hydrogenolysis of)

RN 82586-53-6 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

IT 82586-55-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

RN 82586-55-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

L12 ANSWER 13 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1983:34598 HCAPLUS

DOCUMENT NUMBER:

98:34598

TITLE:

Substituted acyl derivatives of

1,2,3,4-tetrahydroisoquinoline-3-carboxylic

acids

INVENTOR(S):

Hoefle, Milton L.; Klutchko, Sylvester

PATENT ASSIGNEE(S):

Warner-Lambert Co. , USA $\,$

SOURCE:

U.S., 8 pp. Cont.-in-part of U.S. Ser. No.

193,767, abandoned.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE: .

English

2

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.	KIND	DATE	APPLICATION NO.	DATE
US 434		Α		US 1981-236397	19810220
ZA 8100	6332	A	19820929	ZA 1981-6332	19810911
IL 6380	06	A1	19850228	IL 1981-63806	19810911
AU 8175	5416	A1 .	19820506	AU 1981-75416	19810917
AU 5512	239	B2	19860424		
FI 8103	I 8103033 A		19820404	FI 1981-3033	19810930
FI 7869	90	В	19890531		
FI 7869	90		19890911		
DK 8104	4360	Α	19820404	DK 1981-4360	19811001
DK 163			19920120		
			19920615		
EP 4960	05	Al	19820414	EP 1981-304541	19811001
EP 4960	05	B1	19870318		
				LU, NL, SE	
				JP 1981-154900	19811001
			19910305		
				EP 1983-102092	19811001
			19840912		
EP 9615		B1			
				LI, LU, NL, SE	
AT 259	74	E	19870415	AT 1981-304541	19811001
AT 2612	20	E	19870415	AT 1983-102092	19811001

	NO NO	8103359 159017		A B	19820405 19880815		NO	1981-3359	19811002
	NO	159017		C	19881123				
		505960		A1	19821216		E.C	1981-505960	19811002
		201787		A5.	19830810		DD	1981-233850	19811002
	DD			-					
		25890		0	19830829		HÜ	1981-2864	19811002
	но	183602	-	В	19840528				
	SU	1148560		A3	19850330			1981-3340548	19811002
	US	4532342		A	19850730		US	1982-386375	19820608
	AU	8652991		A1	19860717		ΑU	1986-52991	19860204
	AU	563683		B2	19870716				
	FI	8801985		A	19880427		FI	1988-1985	19880427
	FI	79839		В	19891130				
	FI	79839		С	19900312		•		
	CA	1331614		A1	19940823		CA	1992-616411	19920623
	CA	1331615		A1	19940823		CA	1992-616412	19920623
	PRIORITY	APPLN.	INFO.:			US	198	30-193767	19801003
						US		31-236397	19810220
			•			CA		31-387002	19810930
			•			FI		31-3033	19810930
						EP		31-304541	19811001
	OMITED CO	NUDGE (C)		CAG	SREACT 98:3	34598	130	11-204241	19011001
OTHER SOURCE(S):			CAS	ONDAUL JOS.	コサンプロ				

OTHER SOURCE (S):

GΙ

RN

(CH₂)
$$_{\rm n}$$
CH (CO₂R¹) NHCHR²CON

R

R

R

R

1

AΒ Antihypertensive acyltetrahydroisquinolinecarboxylates I (R = H, F, Cl, Br, alkyl, alkoxy, HO, H2N; R1 = H, alkyl; R2 = H, alkyl, PhCH2; R3 = H, alkyl, phenylalkyl; R4, R5 = H, alkyl, alkoxy, alkylthio, alkylsulfonyl, HO; R4R5 = OCH2O; n = 0-3) were prepd. Thus, substitution reaction of H-Ala-OCMe3 with Ph(CH2)2CHBrCO2Et and subsequent debutylation and sepn. of diastereoisomers gave (S,S)-PhCH2CH2CH(CO2Et)-Ala-CO2H, which was condensed with benzyl (S)-1,2,3,4-tetrahydro-6,7-dimethoxy-3-isoquinolinecarboxylate to give (S,S,S)-I (R = H, R1 = Et, R2 = Me, R3 = PhCH2, R4 = R5 = MeO).I inhibited angiotensin converting enzyme with IC50 2.8 .times. 10-9-2.0 .times. 10-6 M.

ΙT 82586-52-5P 82586-55-8P 82637-58-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and angiotensin-converting enzyme inhibiting activity of)

82586-52-5 HCAPLUS CN -3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, monohydrochloride, (3S) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 82586-55-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

HCl

RN 82637-58-9 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-,phenylmethyl ester, (3S)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 82637-57-8 CMF C34 H40 N2 O7

Absolute stereochemistry.

CRN 110-16-7 CMF C4 H4 O4

Double bond geometry as shown.

IT 84048-96-4P

RN 84048-96-4 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, monohydrochloride, [3S-[2[R*(R*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

HCl

IT 82637-57-8P

RN 82637-57-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-,phenylmethyl ester, [3S-[2[R*(R*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L12 ANSWER 14 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1982:492759 HCAPLUS

DOCUMENT NUMBER:

97:92759

TITLE:

Amino acid derivatives, compositions containing

them and their use

INVENTOR(S):

Geiger, Rolf; Teetz, Volker; Urbach, Hansjoerg;

Schoelkens, Bernward; Henning, Rainer

PATENT ASSIGNEE(S):

Hoechst A.-G. , Fed. Rep. Ger.

09/424673

SOURCE: Eur. Pat. Appl., 196 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	PATENT NO.		KIND DATE		AP	DATE	
EP	46953 46953 46953			19820310 19820505 19891206	EP	1981-106535	19810822
	R: AT, 3032709 3118191	BE,	A1	FR, GB, 19820429 19821125	DE	NL, SE 1980-3032709 1981-3118191	
EP	278530 278530	D D	A2 A3	19880817 19890802	EP	1988-102408	19810822
	328160 328160		A1 B1	19890816 19940504	EP	LU, NL, SE 1989-105371	19810822
	R: AT, 48415 105301	BE,	CH, DE, E E	FR, GB, 19891215 19940515	ΑT	LU, NL, SE 1981-106535 1989-105371	19810822 19810822
FI	504955 8102652 90072		Α	19820816 19820301 19930915		1981-504955 1981-2652	
FI HU	90072 27874 189531	•	С	19931227 19831128 19860728	ни	1981-2478	19810827
DK DK	8103835 169382		A B1	19820301 19941017		1981-3835	19810828
AU	8102933 8174718 544756		A A1 B2	19820301 19820311 19850613	AU	1981-2933 1981-74718	
IL	8105988 63683 01048918		A A1 B4	19820825 19880331 19891020	${ t IL}$	1981-5988 1981-63683 1981-134401	19810828 19810828 19810828
ES ES	505604 505605 5158959		A1 A1 A	19821116 19821116 19921027	ES ES	1981-505604 1981-505605 1983-565900	19810918 19810918 19831227
US AU	5162362 8779284		A Al	19921110 19880204	US	1983-565887 1987-79284	19831227 19831227 19871001
JP JP	599151 01125398 06078355		B2 A2 B4	19900712 19890517 19941005		1988-209625	19880825
AU	8936625 627741 04217994			19891005 19920903 19920807		1989-36625 1991-77208	19890620 19910318
JP FI	07121955 90069 90069			19951225 19930915 19931227	FI	1991-4555	19910927
FI FI	90532 90532		B C	19931115 19940225		1991-4554	19910927
US PRIORITY	5401766 APPLN. 1	INFO.	A :	19950328	DE 19 DE 19 EP 19	1994-208443 80-3032709 81-3118191 81-106535 89-105371	19940309 19800830 19810508 19810822 19810822

US 1981-297191

19810828

OTHER SOURCE(S):

CASREACT 97:92759

GΙ

$$CO_2H$$
 CO_2R^3
 CO_2R^3
 CO_2R^5
 CO_2R^5

AΒ Amino acid derivs. I (X = fused benzene or cyclohexane ring; R, R1 =alkyl, alkenyl, cycloalkyl, cycloalkenyl, cycloalkylalkyl, aryl, partially hydrogenated aryl, aralkyl, heterocyclic residue; R2 = H, alkyl, alkenyl, aralkyl; n= 0, 1) were prepd. as long-lasting antihypertensives (no data). Thus, tetrahydroisoquinoline II (R3 = R4 = H) was treated with ZCl (Z = PhCH2O2C) to give II (R3 = H, R4 = Z), which was esterified with Me3COH by DCC in CH2Cl2 contg. 4-(dimethylamino) pyridine to give 97% II (R3 = CMe3, R4 = Z), which was Z-deblocked by hydrogenolysis and then condensed with Z-Ala-OH by DCC/1-hydroxybenzotriazole to give II (R3 = CMe3, R4 = Z-Ala). The latter was Z-deblocked by hydrogenolysis to give II (R = CMe3, R4 = Ala), which condensed with PhCH2CH2COCO2H and was then reduced with NaBH3CN to give isoquinoline III (R5 = CMe3), which was debutylated by CF3CO2H to give III (R5 = H).

82713-51-7P 82713-52-8P 82713-55-1P ΙT 82713-58-4P 82713-62-0P 82713-63-1P 82713-64-2P 82713-65-3P 82713-66-4P 82713-90-4P 82714-28-1P 82714-29-2P 82714-32-7P 82768-84-1P RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 82713-51-7 HCAPLUS

3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-(2-CN methylphenyl)propyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro- (9CI) (CA INDEX NAME)

RN 82713-52-8 HCAPLUS

3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-CN

phenylpropyl]amino]-3-hydroxy-1-oxopropyl]-1,2,3,4-tetrahydro- (9CI)
 (CA INDEX NAME)

RN 82713-55-1 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[3-(4-chlorophenyl)-1-(ethoxycarbonyl)propyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI)(CA INDEX NAME)

RN 82713-58-4 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[6-[(aminoiminomethyl)amino]-2-[[3-(4-chlorophenyl)-1-(ethoxycarbonyl)propyl]amino]-1-oxohexyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82713-62-0 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-(4-fluorophenyl)propyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82713-63-1 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[3-(3,4-dimethoxyphenyl)-1-(ethoxycarbonyl)propyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI)(CA INDEX NAME)

RN 82713-64-2 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[3-(5-chloro-2-methoxyphenyl)-1-(ethoxycarbonyl)propyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82713-65-3 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[3-(2,6-dichlorophenyl)-1-(ethoxycarbonyl)propyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82713-66-4 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-(3,4,5-trimethoxyphenyl)propyl]amino]-3-(1H-imidazol-4-yl)-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82713-90-4 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[3-(5-chloro-2-methoxyphenyl)-1-(ethoxycarbonyl)propyl]amino]-4-methyl-1-oxopentyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82714-28-1 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[2-ethoxy-1-[(3-methoxyphenyl)methyl]-2-oxoethyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82714-29-2 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-[(4-chlorophenyl)methyl]-2-ethoxy-2-oxoethyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82714-32-7 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[2-ethoxy-1-[(4-methylphenyl)methyl]-2-oxoethyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

RN 82768-84-1 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-(9CI) (CA INDEX NAME)

L12 ANSWER 15 OF 15 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1982:472257 HCAPLUS

DOCUMENT NUMBER:

97:72257

TITLE:

Substituted acyl derivatives of

1,2,3,4-tetrahydroisoquinoline-3-carboxylic

acids, their salts and pharmaceutical

compositions containing the derivatives or salts

INVENTOR(S):

Hoefle, Milton Louis; Klutchko, Sylvester

PATENT ASSIGNEE(S):

Warner-Lambert Co. , USA

SOURCE:

Eur. Pat. Appl., 32 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English .

2

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

. Е	PATENT NO.				KIND DATE				I	APP	DATE			
	EP 4960	-		A.		1982			I	EP	1981-3	04541	198110	01
E	EP 4960 R:	5 AT,	BE,	B: CH,		1987 FR,		IT,	LU	, N	IL, SE			
Ţ	JS 4344	949	·	À	•	1982	0817	•	Ţ	US	1981-2	36397	198102	20
E	EP 9615	7		A.	2	1983	1221		I	EΡ	1983-1	02092	198110	01
E	EP 9615	7		A.	3	1984	0912							
E	EP 9615	7		B.	1	1987	0325							
	R:	ΑT,	BE,	CH,	DE,	FR,	GB,	ΙT,	LI	, L	U, NL,	SE		
P	AT 2597	4		E		1987	0415		1	TA	1981-3	04541	198110	01
P	AT 2612	0		E		1987	0415		I	TΑ	1983-1	02092	198110	01
PRIORI	TY APP	LN.	INFO	. :					US :	198	0-1937	67	198010	03
									US :	198	1-2363	97	198102	20
									EP :	198	1-3045	41	198110	01
~-														

GI

$$\begin{array}{c} \text{R}^{3}\text{O}_{2}\text{C} \\ \text{R}\left(\text{CH}_{2}\right)_{n}\text{CH}\left(\text{CO}_{2}\text{R}^{1}\right)\text{NHCHR}^{2}\text{CON} \end{array}$$

Isoquinolinecarboxylates I (R = Ph, substituted Ph; R1 = H, alkyl; R2 = H, alkyl, CH2Ph; R3 = H, alkyl, aralkyl; R4, R5 = H, alkyl, alkoxy, alkylthio, alkylsulfinyl, alkylsulfonyl, OH; R4R5 = OCH2O; n = 0-3) were prepd. Thus, 6.7-dimethoxy-1, 2.3.4-tetrahydro-3-isoquinolinecarboxylic acid was converted to its benzyl ester and treated with (S,S)-HO2CCHMeNHCH(CO2Et)CH2CH2Ph (II) to give I (R = Ph, R1 = Et, R2 = Me, R3 = CH2Ph, R4 = R5 = OMe, n = 2) which was hydrogenated to I (R = Ph, R1 = Et, R2 = Me, R3 = H, R4 = R5 = OMe, n = 2) (III). II was obtained by treating H-Ala-OCMe3 with PhCH2CH2CHBrCO2Et and hydrolysis. III had an angiotensin converting enzyme-inhibiting ED50 of 5.6 .times. 10-9 M in vitro.

IT 82586-53-6P 82637-57-8P 82637-58-9P
RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of)

RN 82586-53-6 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 82637-57-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-,phenylmethyl ester, [3S-[2[R*(R*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 82637-58-9 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, phenylmethyl ester, (3S)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 82637-57-8 CMF C34 H40 N2 O7

Absolute stereochemistry.

CM 2

CRN 110-16-7 CMF C4 H4 O4

Double bond geometry as shown.

ΙT 82586-51-4P 82586-54-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn., hydrogenation, and antihypertensive activity of) 82586-51-4 HCAPLUS

RN

3-Isoquinolinecarboxylic acid, 2-[2-[[1-(ethoxycarbonyl)-3-CN phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, phenylmethyl ester, monohydrochloride, [3S-[2[R*(R*)],3R*]]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 82586-54-7 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, phenylmethyl ester, (3S)-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 82586-53-6 CMF C32 H36 N2 O5

Absolute stereochemistry. Rotation (-).

CM 2

CRN 110-16-7 CMF C4 H4 O4

Double bond geometry as shown.

IT 82586-52-5P 82586-55-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn., hydrolysis, and antihypertensive activity of)

RN 82586-52-5 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-6,7-dimethoxy-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

RN 82586-55-8 HCAPLUS

CN 3-Isoquinolinecarboxylic acid, 2-[(2S)-2-[[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydro-, monohydrochloride, (3S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

● HCl

FILE 'REGISTRY' ENTERED AT 12:03:15 ON 21 APR 2003

L13 38 SEA FILE=REGISTRY ABB=ON PLU=ON (82586-55-8/BI OR 82586-53-6/BI OR 85441-61-8/BI OR 82586-52-5/BI OR 82637-57-8/BI OR 82637-58-9/BI OR 82586-54-7/BI OR 82768-84-1/BI OR 89300-89-0/BI OR 103775-05-9/BI OR 103775-06-0/BI OR 103775-09-3/BI OR 103775-10-6/BI OR 103775-11-7/BI OR 103775-12-8/BI OR 103833-14-3/BI OR 103833-15-4/BI OR 103833-16-5/BI OR 143381-51-5/BI OR 82586-51-4/BI OR 82713-51-7/BI OR 82713-52-8/BI OR 82713-55-1/BI OR 82713-58-4/BI OR 82713-62-0/BI OR 82713-63-1/BI OR 82713-64-2/BI OR 82713-65-3/BI OR 82713-66-4/BI OR 82713-90-4/BI OR 82714-28-1/BI OR 82714-29-2/BI OR 82714-32-7/BI OR 84048-96-4/BI OR 85892-46-2/BI OR 89083-69-2/BI OR 89245-28-3/BI OR 89300-87-8/BI) FILE 'CAOLD' ENTERED AT 12:03:32 ON 21 APR 2003 . 0 S L13 L14 FILE 'USPATFULL' ENTERED AT 12:03:40 ON 21 APR 2003 L15 12 S L13/P L15 ANSWER 1 OF 12 USPATFULL ACCESSION NUMBER: 1999:137527 USPATFULL TITLE: Process for preparing N-[1- (S)-ethoxycarbonyl-3phenylpropyl]-L-alanine derivatives Yang, Suh-Wan, Tao-yuan Hsien, Taiwan, Province INVENTOR(S): of China Chang, Yu-An, Tao-yuan Hsien, Taiwan, Province of China Liu, Yu-Liang, Taipei, Taiwan, Province of China Everlight USA, Inc., Pineville, NJ, United States PATENT ASSIGNEE(S): (U.S. corporation) KIND DATE NUMBER _____ US 5977380 19991102 PATENT INFORMATION: US 1999-251341 19990217 (9) APPLICATION INFO.: Utility DOCUMENT TYPE: Granted FILE SEGMENT: PRIMARY EXAMINER: Powers, Fiona T. Bacon & Thomas LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: 8 EXEMPLARY CLAIM: 1 168 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. A process for synthesizing N-[1-(S)-ethoxycarbonyl-3-phenylpropyl]-L-alanine derivatives of the following formula (I): ##STR1## in which the definition of R has the same meaning as given in the description by using a sulfite derivative to activate the C-terminus of the three dimensional structure of an amino acid of N-[1-(S)-ethoxycarbonyl-3- phenylpropyl]-L-alanine, which can

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 2 OF 12 USPATFULL

ACCESSION NUMBER: 1999:19330 USPATFULL

Searcher: Shears 308-4994

effectively couple with another amino acid to form a dipeptide of formula (I). The compound of fomula (I) is an inhibitor of ACE.

TITLE: Process for preparing an angiotensin converting

enzyme inhibitor

INVENTOR(S): Wang, Shin-Shin, Hsinchu, Taiwan, Province of

China

Tsai, Hui-Ping, Changhua, Taiwan, Province of

China

PATENT ASSIGNEE(S): Industrial Technology Research Institute,

Hsinchu, Taiwan, Province of China (non-U.S.

corporation)

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Barts, Samuel

LEGAL REPRESENTATIVE: Sughrue, Mion, Zinn, Macpeak & Seas, PLLC

NUMBER OF CLAIMS: 18 EXEMPLARY CLAIM: 1 LINE COUNT: 290

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A process for preparing an angiotensin converting enzyme

inhibitor. Phosphorus pentachloride is reacted with the carboxylic

acid group of an amino acid to form an acyl chloride

hydrochloride. The resulting hydrochloride salt is then coupled with a silylated amino acid in a non-aqueous medium to form a high yield peptide. The peptide is used as an angiotensin converting enzyme (ACE) inhibitor.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 3 OF 12 USPATFULL

ACCESSION NUMBER: 95:27333 USPATFULL

TITLE: Aminoacid derivatives, a process for their

preparation, agents containing these compounds,

and the use thereof

INVENTOR(S): Geiger, Rolf, Frankfurt am Main, Germany, Federal

Republic of

Teetz, Volker, Hofheim am Taunus, Germany,

Federal Republic of

Urbach, Hansjorg, Kronberg/Taunus, Germany,

Federal Republic of

Scholkens, Bernward, Kelkheim, Germany, Federal

Republic of

Henning, Rainer, Giessen, Germany, Federal

Republic of

PATENT ASSIGNEE(S): Hoechst Aktiengesellschaft, Frankfurt am Main,

Germany, Federal Republic of (non-U.S.

corporation)

RELATED APPLN. INFO.: Continuation of Ser. No. US 1981-297191, filed on

28 Aug 1981, now abandoned

DATE NUMBER -----PRIORITY INFORMATION: DE 1980-30327094 19800830 19810508 DE 1981-31181910 DOCUMENT TYPE: Utility Granted FILE SEGMENT: PRIMARY EXAMINER: Lone, Werren B. Curtis, Morris & Safford LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: 13 EXEMPLARY CLAIM: 1 2404 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. Aminoacid derivatives of the formula I ##STR1## A , n, R.sup.1, AB R.sup.2 and R.sup.3 have the meanings given, and salts thereof, a process for their preparation, pharmaceutical preparations based on these compounds, and their use as medicaments. CAS INDEXING IS AVAILABLE FOR THIS PATENT. L15 ANSWER 4 OF 12 USPATFULL 92:97073 USPATFULL ACCESSION NUMBER: TITLE: Octahydroindole-2-carboxylic acids Geiger, Rolf, Frankfurt am Main, Germany, Federal INVENTOR(S): Republic of Teetz, Volker, Hofheim am Taunus, Germany, Federal Republic of Urbach, Hansjorg, Kronberg/Taunus, Germany, Federal Republic of Scholkens, Bernward, Kelkheim, Germany, Federal Republic of Henning, Rainer, Giessen, Germany, Federal Republic of Hoechst Aktiengesellschaft, Frankfurt am Main, PATENT ASSIGNEE(S): Germany, Federal Republic of (non-U.S. corporation) NUMBER KIND DATE _____ PATENT INFORMATION: US 5162362 19921110 US 1983-565887 APPLICATION INFO.: 19831227 (6) Division of Ser. No. US 1981-297191, filed on 28 RELATED APPLN. INFO.: Aug 1981 NUMBER DATE DE 1980-3032709 19800830 PRIORITY INFORMATION: 19810508 DE 1981-3118191 Utility DOCUMENT TYPE: FILE SEGMENT: Granted PRIMARY EXAMINER: Vaughn, V. ASSISTANT EXAMINER: Turnipseed, James H. LEGAL REPRESENTATIVE: Curtis, Morris & Safford NUMBER OF CLAIMS: 8 EXEMPLARY CLAIM: 1 LINE COUNT: 1372 CAS INDEXING IS AVAILABLE FOR THIS PATENT. AB What are disclosed are aminoacid compounds of the formula ##STR1## wherein n is 0 or 1, and salts thereof, said compounds and salts

having hypotensive properties; methods for making the compounds; pharmaceutical compositions containing the compounds or salts; and use of the compounds and salts for treating hypertension.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 5 OF 12 USPATFULL

92:89070 USPATFULL ACCESSION NUMBER:

TITLE: Decahydroisoquinoline carboxylic acids

Geiger, Rolf, Frankfurt am Main, Germany, Federal INVENTOR(S):

Republic of

Teetz, Volker, Hofheim am Taunus, Germany,

Federal Republic of

Urbach, Hansjorg, Kronberg/Taunus, Germany,

Federal Republic of

Scholkens, Bernward, Kelkheim (Taunus), Germany,

Federal Republic of

Henning, Rainer, Giessen, Germany, Federal

Republic of

Hoechst Aktiengesellschaft, Frankfurt am Main, PATENT ASSIGNEE(S):

Germany, Federal Republic of (non-U.S.

corporation)

NUMBER KIND DATE _____ US 5158959 19921027 US 1983-565900 19831227 (6) PATENT INFORMATION:

APPLICATION INFO.:

Division of Ser. No. US 1981-297191, filed on 28 RELATED APPLN. INFO.:

Aug 1981

NUMBER DATE _____ DE 1980-3032709 19800830 DE 1981-3118191 19810508 PRIORITY INFORMATION:

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Ivy, C. Warren ASSISTANT EXAMINER: Turnipseed, James H. LEGAL REPRESENTATIVE: Curtis, Morris & Safford

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1525 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

What are disclosed are aminoacid compounds of the formula ##STR1## wherein n is 0 or 1, and salts thereof, said compounds and salts having hypotensive properties; methods for making the compounds; pharmaceutical compositions containing the compounds or salts; and use of the compounds and salts for treating hypertension.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 6 OF 12 USPATFULL

91:96454 USPATFULL ACCESSION NUMBER:

Process for the preparation of n octahydropenta TITLE:

(6) pyrrole carboxylates

Teetz, Volker, Hofheim am Taunus, Germany, INVENTOR(S):

Federal Republic of

Wissmann, Hans, Bad Soden am Taunus, Germany,

Federal Republic of

Urbach, Hansjorg, Kronberg/Taunus, Germany,

Federal Republic of

PATENT ASSIGNEE(S):

Hoechst Aktiengesellschaft, Frankfurt am Main,

Germany, Federal Republic of (non-U.S.

corporation)

KIND DATE NUMBER _____ US 5068351 19911126 US 1990-560004 19900727 (7)

PATENT INFORMATION: APPLICATION INFO.:

RELATED APPLN. INFO.:

Continuation of Ser. No. US 1989-403919, filed on 7 Sep 1989, now abandoned which is a continuation of Ser. No. US 1988-178767, filed on 30 Mar 1988, now abandoned which is a continuation of Ser. No. US 1986-943882, filed on 19 Dec 1986, now abandoned which is a continuation of Ser. No. US 1984-650715, filed on 14 Sep 1984, now abandoned

> DATE NUMBER

PRIORITY INFORMATION:

DE 1983-3333454

19830916

DOCUMENT TYPE: FILE SEGMENT:

Utility Granted

PRIMARY EXAMINER:

Lee, Lester L.

LEGAL REPRESENTATIVE:

Finnegan, Henderson, Farabow, Garrett & Dunner

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

18 1

LINE COUNT:

850

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention relates to a process for the preparation of AB compounds of the formula I ##STR1## in which n is 1 or 2, R denotes hydrogen or an organic radical, R.sup.1 denotes an organic radical, R.sup.2 and R.sup.3 are identical or different and denote hydrogen or an organic radical, and R.sup.4 and R.sup.5, together with the atoms bearing them, form a monocyclic, bicyclic or tricyclic heterocyclic ring system having 5 to 15 carbon atoms, which process comprises reacting compounds of the formula II defined in the description with compounds of the formula IV defined in the description, in the presence of phosphinic anhydrides of the formula III, where appropriate eliminating radicals which have been introduced to protect other functional groups and, where appropriate, esterifying free carboxyl groups in a manner known per se.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 7 OF 12 USPATFULL

ACCESSION NUMBER:

INVENTOR(S):

91:82338 USPATFULL

TITLE:

Process for the preparation of

octahydropenta(b)pyrrole carboxylates Urbach, Hansjorg, Kronberg/Taunus, Germany,

Federal Republic of

Henning, Rainer, Hattersheim am Main, Germany,

Federal Republic of

Wissmann, Hans, Bad Soden am Taunus, Germany,

Federal Republic of

Teetz, Volker, Hofheim am Taunus, Germany,

308-4994 Searcher : Shears

Federal Republic of

PATENT ASSIGNEE(S): Hoechst Aktiengesellschaft, Frankfurt am Main,

Germany, Federal Republic of (non-U.S.

corporation)

NUMBER KIND DATE

US 5055591 19911008
US 1988-173024 19880323 (7)

APPLICATION INFO.: US 1988-173024
RELATED APPLN. INFO.: Continuation of Ser. N

Continuation of Ser. No. US 1986-943881, filed on

19 Dec 1986, now abandoned which is a

continuation of Ser. No. US 1984-650714, filed on

14 Sep 1984, now abandoned

PRIORITY INFORMATION:
DOCUMENT TYPE:
FILE SEGMENT:

PATENT INFORMATION:

Utility Granted

PRIMARY EXAMINER: Lee,

Lee, Lester L.

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM: 1

LEGAL REPRESENTATIVE: Finnegan, Henderson, Farabow, Garrett, and Dunner NUMBER OF CLAIMS: 20

LINE COUNT: 855

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention relates to a process for the preparation of compounds of the formula I ##STR1## in which n is 1 or 2, R denotes hydrogen or an organic radical, R.sup.1 denotes an organic radical, R.sup.2 and R.sup.3 are identical or different and denote hydrogen or an organic radical, and R.sup.4 and R.sup.5, together with the atoms bearing them, form a monocyclic, bicyclic or tricyclic heterocyclic ring system having 3 to 15 carbon atoms, which process comprises reacting compounds of the formula II defined in the description with compounds of the formula III defined in the description, in the presence of alkanephosphonic anhydrides, where appropriate eliminating radicals which have been introduced to protect other functional groups and, where appropriate, esterifying free carboxyl groups in a manner known per se.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 8 OF 12 USPATFULL

ACCESSION NUMBER: 88:48826 USPATFULL

TITLE: Crystalline quinapril and a process for producing

the same

INVENTOR(S): Goel, Om P., Ann Arbor, MI, United States

Krolls, Uldis, Ann Arbor, MI, United States

PATENT ASSIGNEE(S): Warner-Lambert Company, Morris Plains, NJ, United

States (U.S. corporation)

NUMBER KIND DATE

-----PATENT INFORMATION: US 4761479 19880802
APPLICATION INFO.: US 1987-32209 19870330 (7)
DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Hollrah, Glennon H.

ASSISTANT EXAMINER: Turnipseed, James H. LEGAL REPRESENTATIVE: Anderson, Elizabeth M.

NUMBER OF CLAIMS: 8
EXEMPLARY CLAIM: 1
LINE COUNT: 237

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A novel crystalline form of quinapril and a novel process for the large scale preparation of the ACE inhibitor, quinapril, in a highly pure state. The substance is of high bulk density suitable for formulation in capsules and tablets. This inexpensive process uses HCl gas in glacial acetic acid for rapid and clean

de-t-butylation at room temperature.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 9 OF 12 USPATFULL

ACCESSION NUMBER: 85:44762 USPATFULL

TITLE: N-substituted amino acids as intermediates in the

preparation of acyl derivatives of

1,2,3,4-tetrahydroisoquinoline-3-carboxylic acids INVENTOR(S): Hoefle, Milton L., Ann Arbor, MI, United States

Klutchko, Sylvestor, Ann Arbor, MI, United States

PATENT ASSIGNEE(S): Warner-Lambert Company, Morris Plains, NJ, United

States (U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 4532342 19850730

APPLICATION INFO.: US 1982-386375 19820608 (6)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1981-236397, filed on 20 Feb 1981, now patented, Pat. No. US 4344949 which is a continuation-in-part of Ser.

No. US 1980-193767, filed on 3 Oct 1980, now abandoned

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted

PRIMARY EXAMINER: Hollrah, Glennon H. ASSISTANT EXAMINER: Turnipseed, James H. LEGAL REPRESENTATIVE: Daignault, Ronald A.

NUMBER OF CLAIMS: 3
EXEMPLARY CLAIM: 1
LINE COUNT: 555

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

N-substituted amino acids are described which when coupled with 1,2,3,4-tetrahydroisoquinolines result in substituted acyl derivatives of 1,2,3,4-tetrahydroisoquinoline-3-carboxylic acids as anti-hypertensive agents. The novel intermediates are in turn prepared by reacting an amino acid such as alanine with 2-bromo-4-phenyl butanoic acid or an ester thereof.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 10 OF 12 USPATFULL

ACCESSION NUMBER: 84:8849 USPATFULL TITLE: Antihypertensive agents

INVENTOR(S): Smith, Elizabeth M., Verona, NJ, United States

Witkowski, Joseph T., Morris Township, Morris

County, NJ, United States

Schering Corporation, Kenilworth, NJ, United PATENT ASSIGNEE(S):

States (U.S. corporation)

NUMBER KIND DATE US 4431645 US 1982-355639 19840214 PATENT INFORMATION: 19820308 (6) APPLICATION INFO.: Utility DOCUMENT TYPE:

Granted FILE SEGMENT: PRIMARY EXAMINER: Ford, John M.

LEGAL REPRESENTATIVE: Magatti, Anita W., Eisen, Bruce M.

17 NUMBER OF CLAIMS: 1,16 EXEMPLARY CLAIM: LINE COUNT: 444

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compounds of the formula ##STR1## and the pharmaceutically acceptable salts thereof, wherein R.sup.1, R.sup.2, R.sup.3 and R.sup.4 are independently selected from hydrogen or lower alkyl; n is 1 or 0; A and B taken together with the carbons to which they are attached form an alkylene ring having six carbon atoms or A and B are hydrogen; and Z is ##STR2## The compounds are useful as hypertensive agents.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 11 OF 12 USPATFULL

PATENT ASSIGNEE(S):

ACCESSION NUMBER: 84:8848 USPATFULL Antihypertensive agents TITLE:

INVENTOR(S):

Smith, Elizabeth M., Verona, NJ, United States Witkowski, Joseph T., Morris Township, Morris

County, NJ, United States

Doll, Ronald J., Maplewood, NJ, United States Schering Corporation, Kenilworth, NJ, United

States (U.S. corporation)

NUMBER KIND DATE -----US 4431644 19840214 US 1982-355638 19820308 (6) PATENT INFORMATION: APPLICATION INFO.: DOCUMENT TYPE: Utility PRIMARY EXAMINER: LEGAL REPRESENTE Granted Ford, John M. Magatti, Anita W., Eisen, Bruce M. LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: 12

EXEMPLARY CLAIM: 1,11

497 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compounds of the formula ##STR1## and the pharmaceutically acceptable salts thereof, wherein R.sup.1, R.sup.2 and R.sup.4 are independently selected from hydrogen and lower alkyl; R.sup.3 is hydrogen, lower alkyl or amino lower alkyl; A and B taken together with the carbons to which they are attached form an alkylene ring having six carbon atoms or A and B are hydrogen; and Z is ##STR2## are disclosed. The compounds are useful as anti-hypertensive agents.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Searcher : Shears

L15 ANSWER 12 OF 12 USPATFULL

ACCESSION NUMBER: 82:39913 USPATFULL

TITLE: Substituted acyl derivatives of

1,2,3,4-tetrahydroisoquinoline-3-carboxylic acids INVENTOR(S): Hoefle, Milton L., Ann Arbor, MI, United States

Klutchko, Sylvester, Ann Arbor, MI, United States

PATENT ASSIGNEE(S): Warner-Lambert Company, Morris Plains, NJ, United

States (U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 4344949 19820817 APPLICATION INFO .: US 1981-236397 19810220 (6)

Continuation-in-part of Ser. No. US 1980-193767, RELATED APPLN. INFO.:

filed on 3 Oct 1980, now abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Daus, Donald G.

Turnipseed, James H. ASSISTANT EXAMINER:

Patton, Walter LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 15 EXEMPLARY CLAIM: 1,15 LINE COUNT: 606

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Substituted acyl derivatives of 1,2,3,4-tetrahydroisoquinoline-3-AR carboxylic acids and the pharmaceutically acceptable salts thereof are produced by coupling a suitably substituted 1,2,3,4-tetrahydroisoguinoline with a suitably substituted amino acid and when desired hydrolyzing or removing protecting groups of the resulting product. The compounds of the invention, their salts and pharmaceutical compositions thereof are useful as antihypertensive agents.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

(FILE 'MARPAT' ENTERED AT 12:04:06 ON 21 APR 2003) L16 STR

REP G1=(1-3) CH2 VAR G2=OH/30

NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

MLEVEL IS CLASS AT 22 26 32

GGCAT IS UNS AT 32

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 32

STEREO ATTRIBUTES: NONE

ATTRIBUTES SPECIFIED AT SEARCH-TIME:

ECLEVEL IS LIM ON ALL NODES ALL RING(S) ARE ISOLATED

L18 17 SEA FILE=MARPAT SSS FUL L16 (MODIFIED ATTRIBUTES)

100.0% PROCESSED 2950 ITERATIONS

17 ANSWERS

DATE

SEARCH TIME: 00.00.20

L18 ANSWER 1 OF 17 MARPAT COPYRIGHT 2003 ACS

ACCESSION NUMBER: 134:157567 MARPAT

TITLE: Method of using angiotensin converting enzyme

inhibitors to stimulate angiogenesis

APPLICATION NO.

INVENTOR(S):
Isner, Jeffrey Michael

PATENT ASSIGNEE(S): Warner-Lambert Company, USA

KIND

SOURCE:

U.S., 21 pp.

CODEN: USXXAM

DATE

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.

US 6191144	B1 200102	20 US	1999-361351	19990726
PRIORITY APPLN. INFO.:	:	US	1998-96814P	19980817
AB The invention is				
stimulate angioge	enesis in man	mals or in	mammalian tiss	ue in vitro.
Specifically, the				
angiogenesis thro				
and to ACE inhib				
enhancement of ar				
in the promotion				
healing, bone hea				
promoting the for				
preferred embodir				
is used to treat,	, prophylacti	cally or ot	herwise, mamma	ls in need of

IC ICM A61K031-445

NCL 514315000

CC 1-8 (Pharmacology)

angiogenic-treatment.

Section cross-reference(s): 63

ST angiotensin converting enzyme inhibitor angiogenesis stimulation; ACE inhibitor angiogenesis stimulation wound healing; bone healing ACE inhibitor angiogenesis stimulation; burn ACE inhibitor angiogenesis stimulation; quinapril quinaprilat angiogenesis

```
stimulation
     Angiogenesis
TT
     Burn
     Drug delivery systems
     Wound healing promoters
        (ACE inhibitors for stimulation of angiogenesis)
ΙT
     Blood vessel
        (collateral; ACE inhibitors for stimulation of angiogenesis)
ΙT
     Bone
        (healing; ACE inhibitors for stimulation of angiogenesis)
IT
     Transplant and Transplantation
        (skin; ACE inhibitors for stimulation of angiogenesis)
IT
        (transplant; ACE inhibitors for stimulation of angiogenesis)
ΙT
     62571-86-2, Captopril 127464-60-2, Vascular endothelial growth
     factor
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); BIOL (Biological study)
        (ACE inhibitors for stimulation of angiogenesis)
     82768-85-2, Quinaprilat 85441-61-8, Quinapril
IT
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (ACE inhibitors for stimulation of angiogenesis)
TΤ
     9015-82-1, Angiotensin converting enzyme
     RL: BPR (Biological process); BSU (Biological study, unclassified);
     BIOL (Biological study); PROC (Process)
        (ACE inhibitors for stimulation of angiogenesis)
                               THERE ARE 24 CITED REFERENCES AVAILABLE
REFERENCE COUNT:
                         24
                               FOR THIS RECORD. ALL CITATIONS AVAILABLE
                               IN THE RE FORMAT
L18 ANSWER 2 OF 17 MARPAT COPYRIGHT 2003 ACS
                         132:15667 MARPAT
ACCESSION NUMBER:
                         Stabilization of formulations containing ACE
TITLE:
                         inhibitors by magnesium oxide
                         Daniel, Jane Ellen; Harris, Michael Ray;
INVENTOR(S):
                         Hokanson, Gerard Clifford; Weiss, Jay
                         Warner-Lambert Company, USA
PATENT ASSIGNEE(S):
SOURCE:
                         PCT Int. Appl., 27 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                      KIND DATE
                                           APPLICATION NO.
                                                             DATE
     PATENT NO.
                      ----
                                           _____
     ______
                                       WO 1999-US10189 19990510
                     A1 . 19991209
     WO 9962560
         W: AE, AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GD, GE, HR,
             HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MG, MK,
             MN, MX, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE,
             DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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CA 1999-2330581 19990510

19990510

AU 1999-39793

CA 2330581

AU ,9939793

AA

Α1

19991209

19991220

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BR 1999-10947
                            20010306
                                                             19990510
     BR 9910947
                       Α
     EP 1083931
                       Α1
                            20010321
                                           EP 1999-922899
                                                             19990510
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
             PT, IE, SI, LT, LV, FI, RO
                                           JP 2000-551814
                                                             19990510
     JP 2002516881
                       Т2
                            20020611
    NZ 508544
                                           NZ 1999-508544
                                                             19990510
                            20021025
                       Α
     US 6417196
                       R1
                            20020709
                                           US 2000-700883
                                                             20001120
     NO 2000006148
                                           NO 2000-6148
                                                             20001204
                       Α
                            20001204
                                                             20020422
     US 2002161020
                       A1
                            20021031
                                           US 2002-127181
                                           US 1998-88280P
                                                             19980605
PRIORITY APPLN. INFO.:
                                           WO 1999-US10189
                                                             19990510
                                           US 2000-700883
                                                             20001120
     The present invention is directed to ACE inhibitor-contg. compns.
AB
     stabilized by the presence of magnesium oxide. Preferably, the ACE
     inhibitor, quinapril, is protected from certain forms of degrdn.
     when prepd. in a pharmaceutical compn. consisting essentially of
    magnesium oxide as the stabilizing agent. The presence of magnesium
     oxide also lends itself to favorable processing conditions during
     the manuf. of ACE inhibitor-contg. compns., esp. processing by wet
     granulation. Thus, tablets contained quinapril-HCl 21.7, MgO 21.7,
     lactose 254.3, gelatin 6.4, Polyplasdone 12.8, and Mg stearate 3.2
          The MgO-contg. formulation required less gelatin than the Mg
     carbonate hydroxide formulation to obtain acceptable compressibility
     and stabilized the formulation.
IC
     ICM A61K047-02
         A61K047-26; A61K031-47
     ICS
CC
     63-6 (Pharmaceuticals)
ST
     stabilization formulation ACE inhibitor magnesium oxide
IT
     Friction
     Stabilizing agents
        (stabilization of formulations contg. ACE inhibitors by magnesium
        oxide)
IT
     Drug delivery systems
        (tablets; stabilization of formulations contg. ACE inhibitors by
        magnesium oxide)
ΤТ
     Granulation
        (wet; stabilization of formulations contg. ACE inhibitors by
        magnesium oxide)
TΤ
     9015-82-1
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitors; stabilization of formulations contg. ACE inhibitors
        by magnesium oxide)
     82586-55-8, Quinapril hydrochloride
                                           85441-61-8, Quinapril
IT
     RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological
     study); USES (Uses)
        (stabilization of formulations contg. ACE inhibitors by magnesium
        oxide)
IΤ
                       69-65-8, Mannitol
                                            1309-48-4, Magnesium oxide
     63-42-3, Lactose
     (MgO), biological studies
                                 75847-73-3, Enalapril
                                                         80876-01-3,
     Indolapril
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (stabilization of formulations contq. ACE inhibitors by magnesium
        oxide)
REFERENCE COUNT:
                               THERE ARE 4 CITED REFERENCES AVAILABLE FOR
                               THIS RECORD. ALL CITATIONS AVAILABLE IN
                               THE RE FORMAT
```

L18 ANSWER 3 OF 17 MARPAT COPYRIGHT 2003 ACS

```
ACCESSION NUMBER:
                         131:322920 MARPAT
                         Process for preparing N-[1(S)-ethoxycarbonyl-3-
TITLE:
                         phenylpropyl]-L-alanine derivatives
INVENTOR(S):
                         Yang, Suh-Wan; Chang, Yu-An; Liu, Yu-Liang
PATENT ASSIGNEE(S):
                         Everlight USA, Inc., USA
SOURCE:
                         U.S., 6 pp.
                         CODEN: USXXAM
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                     KIND
                            DATE
                                          APPLICATION NO.
                                                            DATE
     -----
                     ____
                            -----
                                          -----
                                                           _____
     US 5977380
                            19991102
                                           US 1999-251341
                                                            19990217
PRIORITY APPLN. INFO.:
                                           US 1999-251341
                                                            19990217
                         CASREACT 131:322920
OTHER SOURCE(S):
AB · (S)-EtO2CCH(CH2CH2Ph)-Ala-R (I; R are certain cyclic amino acids,
     e.g., L-proline) or their pharmaceutically acceptable salts were
     prepd. by coupling I (R = OC6H4R1, where R1 is nitro, cyano,
     sulfite, carboxy, aldehyde, ester, or halo) with an amino acid.
     Thus, I (R = OC6H4NO2-p), formed by esterifying the acid with
     4-nitrophenol in the presence of triethylamine and thionyl chloride
     in dichloromethane, was treated with L-proline to afford I (R =
     proline residue) (enalapril).
IC
     ICM C07D207-12
     ICS C07D233-26; C07D217-16; C07D495-10
NCL
     548533000
CC
     34-3 (Amino Acids, Peptides, and Proteins)
ST
     ethoxycarbonylphenylpropylalanyl dipeptide prepn; peptide
     ethoxycarbonylphenylpropylalanyl prepn; enalapril prepn
ΙT
     Dipeptides
     RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of [(ethoxycarbonyl)phenylpropyl]-L-alanine derivs.)
TΤ
     75847-73-3P, Enalapril 76095-16-4P, Enalapril maleate
     80876-01-3P
                  83059-56-7P
                                 83435-66-9P
                                              83647-97-6P
                                                             85441-61-8P
     87333-19-5P
                   89371-37-9P
                                 103775-10-6P
                                               109683-61-6P
     RL: IMF (Industrial manufacture); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of [(ethoxycarbonyl)phenylpropyl]-L-alanine derivs.)
ΙT
     147-85-3, L-Proline, reactions 25887-81-4
                                                   74163-81-8
     80871-70-1
                 80875-98-5
                             82717-96-2
                                          83552-44-7
                                                        103733-66-0
     107716-98-3
                  109428-53-7
                                 109583-12-2
                                               248254-17-3
                                                             248254-18-4
     248254-19-5
                   248606-38-4
                                 248606-39-5
                                               248606-40-8
                                                             248606-41-9
     248606-42-0
                   248606-43-1
                                 248606-44-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of [(ethoxycarbonyl)phenylpropyl]-L-alanine derivs.)
IT
     248254-21-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (prepn. of [(ethoxycarbonyl)phenylpropyl]-L-alanine derivs.)
REFERENCE COUNT:
                         5
                               THERE ARE 5 CITED REFERENCES AVAILABLE FOR
                               THIS RECORD. ALL CITATIONS AVAILABLE IN
                               THE RE FORMAT
L18 ANSWER 4 OF 17 MARPAT COPYRIGHT 2003 ACS
```

Searcher: Shears 308-4994

130:168658 MARPAT

ACCESSION NUMBER:

TITLE: Process for preparing pharmacologically

acceptable salts of N-[1(S)-ethoxycarbonyl-3-

phenylpropyl]-L-alanyl amino acids

INVENTOR(S): Ueda, Yasuyoshi; Kinoshita, Koichi; Moroshima,

Tadashi; Yanagida, Yoshifumi; Fuse, Yoshihide

PATENT ASSIGNEE(S): Kaneka Corporation, Japan SOURCE: PCT Int. Appl., 97 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

LANGUAGE:

GI

PA	PATENT NO.				KIND DATE				APPLICATION NO.						DATE		
WO	7O 9905164			A1 19990204				WO 1998-JP3240						19980721			
	W:	CA,	CN,	HU,	IL,	JP,	KR,	SG,	SI,	US							
	RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	
		NL,	PT,	SE													
EP	9672	21		A.	1	1999	1229		EF	19	98-9	3258	5	1.998	0721		
	R:	AT,	CH,	DE,	ES,	FR,	GB,	ΙΤ,	LI,	NL,	ΙE						
US	6335	453		B.	1	2002	0101		US	19	99-2	6910	7	1999	0319		
US	2002	0870	07	A.	1	2002	0704		US	20	01-9	89186	6	2001	1121		
US	6518	436		· B2	2	2003	0211										
PRIORIT	Y APP	LN.	INFO	. :					JE	19	97-1	9586	5	19970	0722		
									WC	19	98-J	P3240)	19980	0721		
OTHER SO	OURCE	(S):			CAS	REAC'	T 13	0:16	8658								

Claimed is a process for prepq. pharmacol. acceptable salts of ΑB N-[1(S)-ethoxycarbonyl-3-phenylpropyl]-L-alanyl amino acids, comprising the steps of: condensing an amino acid with N-[1(S)-ethoxycarbonyl-3-phenylpropyl]-L-alanine N-carboxy anhydride (I) under basic conditions; decarboxylating the condensate under neutral to acidic conditions to prep. an N-[1(S)-ethoxycarbonyl-3phenylpropyl]-L-alanyl amino acid; and converting the product to a pharmacol. acceptable salt thereof, characterized in that a series of procedures up to the formation of a pharmacol. acceptable salt or up to the withdrawal of the pharmaceutically acceptable salt thereof are carried out in an aq. liq. to inhibit the prodn. of a byproduct diketopiperazine, e.g. II. According to this process, high-quality pharmacol. acceptable salts of N-[1(S)-ethoxycarbonyl-3phenylpropyl]-L-alanyl amino acids can be prepd. in high yields in a cost-effective manner on a com. scale. Thus, a soln. of 29.20 g I in 156 mL EtOAc was slowly added dropwise over 4 h at 19-20.degree.

```
to a mixt. of 22.02 g L-proline, 20 mL EtOAc, and 22 mL H2O
     (adjusted to pH 10.5 by adding 30 wt.% aq. NaOH) with stirring,
    while the pH of the reaction mixt. was kept at pH 10.5.+-.0.5 by
    adding 30 wt.% aq. NaOH during the reaction. After completing the
    addn., the reaction mixt. was stirred for another 1 h under the same
    condition, warmed to 30.degree., made pH 4.5.+-.0.2 by adding 35%
    wt.% aq. HCl, and stirred for 10 min to complete decarboxylation.
    The org. phase was sepd. and the aq. phase was extd. once with
    EtOAc. The ext. was combined and washed once with 5% vol. of H2O to
    qive the water-satd. org. phase contg. N-[1(S)-ethoxycarbonyl-3-
    phenylpropyl]-L-alanyl-L-proline (enalapril) (III) 14, II 0.5,
    N-[1(S)-carboxy-3-phenylpropyl]-L-alanyl-L-proline 0.4, and
    N-[1(S)-ethoxycarbonyl-3-phenylpropyl]-L-alanine 0.5 wt.%. To the
    org. phase was added 10.49 g maleic acid and the resulting mixt. was
    stirred at 30.degree. for 1 h, cooled to 3.degree. over 3 h, and was
    stirred for another 2 h to give, after filtration of the pptd.
    crystals, washing them with EtOAc chilled to 5.degree., and vacuum
    drying, 90% III maleate of .gtoreq.99% purity.
     ICM C07K005-062
    ICS C07K001-02; C07K001-08
     34-3 (Amino Acids, Peptides, and Proteins)
    Section cross-reference(s): 1
    ethoxycarbonylphenylpropylalanyl amino acid prepn enalapril;
    ethoxycarbonylphenylpropylalanine carboxy anhydride condensation
    amino acid
    Dipeptides
    RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of pharmacol. acceptable salts of N-[1(S)-ethoxycarbonyl-
        3-phenylpropyl]-L-alanyl amino acids)
    76420-72-9P
                   82717-96-2P
                                 115729-52-7P
    RL: BYP (Byproduct); REM (Removal or disposal); PREP (Preparation);
    PROC (Process)
        (prepn. of pharmacol. acceptable salts of N-[1(S)-ethoxycarbonyl-
        3-phenylpropyl]-L-alanyl amino acids)
     147-85-3, L-Proline, reactions
                                      67123-97-1, 1,2,3,4-
    Tetrahydroisoguinoline-3-carboxylic acid
                                                84793-24-8
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of pharmacol. acceptable salts of N-[1(S)-ethoxycarbonyl-
        3-phenylpropyl]-L-alanyl amino acids)
     220069-65-8P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of pharmacol. acceptable salts of N-[1(S)-ethoxycarbonyl-
        3-phenylpropyl]-L-alanyl amino acids)
                              76095-16-4P, Enalapril maleate
     75847-73-3P, Enalapril
     RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of pharmacol. acceptable salts of N-[1(S)-ethoxycarbonyl-
        3-phenylpropyl]-L-alanyl amino acids)
                               THERE ARE 25 CITED REFERENCES AVAILABLE
REFERENCE COUNT:
                         25
                               FOR THIS RECORD. ALL CITATIONS AVAILABLE
                               IN THE RE FORMAT
                     MARPAT COPYRIGHT 2003 ACS
L18 ANSWER 5 OF 17
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ACCESSION NUMBER:

INVENTOR(S):

TITLE:

308-4994 Searcher : Shears

Wang, Shin-Shin; Tsai, Hui-Ping

Process for preparing enalapril and related

angiotensin converting enzyme inhibitors

130:139658 MARPAT

PATENT ASSIGNEE(S): Industrial Technology Research Institute, Taiwan

SOURCE: U.S., 5 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

COLINIA

	LY ACC. NUM. COUN NT INFORMATION:	T: 1								
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE					
OTHE	US 5869671 RITY APPLN. INFO. R SOURCE(S):	: CA	SREACT 130:	US 1998-106288 139658	19980629					
AB	alkyl; R3 = alky = an amino acid enzyme inhibitor phenylpropyl]-L- with L-proline i 79.4% enalapril.	l, Ph, residu s. Th alanin	phenylmeth e) were pre us, treatin e with PC15	= R302CCHR4NH or AcSC yl; R4 = phenylalkyl, pd. as angiotensin co g N-[1-(S)-ethoxycarb and coupling of the hexamethyldisilazane	Ph, alkyl; A nverting onyl-3- acyl chloride					
IC NCL	ICM C07D217-00 546147000	•								
CC										
ST	enalapril prepn inhibitor angiot			l prepn process; pept enzvme	ide prepn					
IT	Peptides, prepar RL: IMF (Industr	ation ial ma	nufacture);	PREP (Preparation) and related angiotens	in converting					

IT 10416-59-8, BSA

enzyme inhibitors)

RL: RCT (Reactant); RACT (Reactant or reagent) (bis(trimethylsilyl)acetamide; process for prepg. enalapril and related angiotensin converting enzyme inhibitors)

IT 9015-82-1, Angiotensin converting enzyme

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(process for prepg. enalapril and related angiotensin converting enzyme inhibitors)

64838-55-7P IT75847-73-3P, Enalapril 85441-61-8P, Quinapril

RL: IMF (Industrial manufacture); PREP (Preparation) (process for prepg. enalapril and related angiotensin converting enzyme inhibitors)

IT 999-97-3, Hmds 74163-81-8 82717-96-2 148493-16-7

RL: RCT (Reactant); RACT (Reactant or reagent)

(process for prepg. enalapril and related angiotensin converting enzyme inhibitors)

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L18 ANSWER 6 OF 17 MARPAT COPYRIGHT 2003 ACS

ACCESSION NUMBER:

129:28219 MARPAT

TITLE:

Preparation of organic nitrates as

antithrombotics. Del Soldato, Piero

INVENTOR(S): PATENT ASSIGNEE(S):

Nicox S.A., Fr.; Del Soldato, Piero

SOURCE:

PCT Int. Appl., 42 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATEN'	r no.		KI	ND	DATE			A	PPLI	CATI	ON N	ο.	DATE		
	WO 982	21193		 A	 1	1998	0522		W	0 199	 97-Е	 P631	1	1997	1112	
	W													IS,		KP,
														RO,		
				TR,	TT,	UA,	US,	UZ,	VN,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,
	D.		TM	T 0	3.67-7	an	0.5	110	C7 7-7	70 CD	DE	CII	DE	DIA	ЕС.	DT
	RI	W: GH,												CF,		
			GA,								SE,	DE,	ъυ,	Cr,	CG,	CI,
	AU 98		GA,	GN,		1998				ນ 19:	98-5	5519		1997	1112	
	AU 72			В		2001				0 10.	, ,	5515		133,		
	EP 94			Ā		1999			E.	P 19	97-9	5189	0	1997	1112	
	EP 94					2002										
	R		BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	NL,	SE,	PT,	ΙE,
		SI,	LT,	FI,	RO											
	CN 12			A		2000			C	N 199	97-1	8124	5	1997	1112	
	CN 10			В		2002								4000		
	BR 97			A		2000				R 199				1997		
	JP 200		76			2001				P 199				1997		
	RU 219					2002				U 199 T 199				1997 1997		
	AT 220 US 624			E B		2002 2001				S 199				1999		
	KR 200		51			2000			K	R 19	99-7	0422	9	1999		
PRTO	RITY A						0023			г 199				1996		
11110			11110	• •						0 19				1997		
AB	A (XNO								= s	peci:	fied	res	idue		•	
	cardio															
	Thus,															
	3-HOC															
	EtOAc															
	at 10			ттА	ın r	ats	gave	538	ant:	ıtnro	oamc	tic	acti	rarch	, vs	. 118
IC	for en															
10	ICS (61 KO	31-4	1 : A	61 KO	38-0	5						
CC	34-3															
	Section									,						
ST	org n								imol	ol n	itra	te d	eriv	, pre	on	
	antit															
	antihy	yperte	ensiv	e or	g ni	trat	e pr	epn;	pla [.]	telet	t ag	greg	atio	on in	hibi	cor
	org n															
ΙT	Antic															
	Antih															
	Broncl Plate				n in	hihi	+0~0									
		repn.						anti	thro	mhot:	icel					
ĪT	Peptio					4663	us (A11 C I	C111 O		.03)					
	RL: B					ivit	y or	eff	ecto	r, e	xcep	t ad	vers	se);	BSU	
	(Biole															THU
	(Ther	apeuti	c us	e);	BIOL	(Bi	olog.	ical	stu	dy);	PRE	P (P	repa	arati	on);	USES
	(Uses) _											-			
	(p:	repn.	of o	rg.	nitr	ates	as a	anti	thro	mbot:	ics)					
								•								

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207987-09-5P 207987-11-9P
                                                 207987-13-1P
    207987-07-3P
TΤ
    RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
        (prepn. of org. nitrates as antithrombotics)
     927-58-2, 4-Bromobutyryl chloride
                                        26839-77-0
                                                     75847-73-3,
ΙT
                76420-72-9
                            190442-16-1
    Enalapril
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of org. nitrates as antithrombotics)
                                            207987-17-5P
                                                           207987-19-7P
     26839-76-9P, R-Timolol
                            207987-15-3P
ΙT
                                  207987-25-5P
    207987-21-1P
                   207987-23-3P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (prepn. of org. nitrates as antithrombotics)
                              THERE ARE 3 CITED REFERENCES AVAILABLE FOR
                         3
REFERENCE COUNT:
                               THIS RECORD. ALL CITATIONS AVAILABLE IN
                               THE RE FORMAT
L18 ANSWER 7 OF 17 MARPAT COPYRIGHT 2003 ACS
                         128:61791 MARPAT
ACCESSION NUMBER:
                         Method for the production of L-alanine
TITLE:
                         derivatives with an ACE inhibitor effect
                         Palomo Coll, Alberto; Serra Mortes, Sonia
INVENTOR(S):
                         KRKA Tovarna Zdravil D. D., Slovenia
PATENT ASSIGNEE(S):
SOURCE:
                         Ger. Offen., 6 pp.
                         CODEN: GWXXBX
                         Patent
DOCUMENT TYPE:
                         German
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                          APPLICATION NO. DATE
                            DATE
     PATENT NO.
                     KIND
                            -----
                                          _____
     ______
                     ____
     DE 19721290
                     A1
                            19971211
                                          DE 1997-19721290 19970521
                                          SI 1996-169
                                                       19960522
PRIORITY APPLN. INFO.:
     Title compds. R1CH2CH2CH(CO2Et)NHCH(CH3)COR2 [(I): R1 = alkyl, aryl,
     heterocycle; R2 = (un)natural .alpha.-amino acid], and their
     pharmaceutically acceptable salts were prepd. as ACE-inhibitors (no
     data). Thus, (S,S)-I (R1 = Ph; R2 = OH) was reacted with L-proline
     to yield (S,S)-I (R1 = Ph; R2 = L-proline), which was converted to
     its maleate salt.
     C07K005-078; C07D285-12; C07D521-00; C07D207-16; C07D217-26;
IC
     CO7D233-32; CO7D209-42; CO7D495-10; CO7D209-52
     34-2 (Amino Acids, Peptides, and Proteins)
CC
     Section cross-reference(s): 1
     amino acid prepn ACE inhibitor; antihypertensive amino acid prepn;
ST
     alanine proline prepn ACE inhibitor
IT
     Antihypertensives
        (prepn. of L-alanine derivs. with an ACE inhibitor effect)
     Amino acids, preparation
IT
     RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); RACT (Reactant or reagent); USES (Uses)
        (prepn. of L-alanine derivs. with an ACE inhibitor effect)
                  87679-37-6P
                                 200423-23-0P
ΙT
     76420-75-2P
     RL: IMF (Industrial manufacture); SPN (Synthetic preparation); THU
    (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
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(Uses)
        (prepn. of L-alanine derivs. with an ACE inhibitor effect)
                   80876-01-3P
                                83059-56-7P
                                               83435-66-9P
                                                             83647-97-6P
TT:
     75847-73-3P
                   87333-19-5P
                                 89371-37-9P
                                               103775-10-6P
     85441-61-8P
     109683-61-6P
     RL: IMF (Industrial manufacture); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of L-alanine derivs. with an ACE inhibitor effect)
                                    80875-98-5
                                                   82717-96-2
IT
     147-85-3, L-Proline, reactions
     145438-94-4
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of L-alanine derivs. with an ACE inhibitor effect)
IT
     200423-22-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (prepn. of L-alanine derivs. with an ACE inhibitor effect)
L18 ANSWER 8 OF 17 MARPAT COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                         128:35025 MARPAT
TITLE:
                         Process for the preparation of
                         N-(1-alkoxycarbonyl-3-phenylpropyl)amino acid
                         derivatives
                         Ueda, Yasuyoshi; Matsumoto, Akira; Manabe,
INVENTOR(S):
                         Hajime
PATENT ASSIGNEE(S):
                         Kaneka Corporation, Japan; Ueda, Yasuyoshi;
                         Matsumoto, Akira; Manabe, Hajime
                         PCT Int. Appl., 74 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         Japanese
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                     KIND DATE
                                           APPLICATION NO.
                                                           DATE
     ______
                     ____
                           _____
                                           _____
                                                            19970508
                           19971120
                                           WO 1997-JP1543
     WO 9743246
                     A1
         W: CA, CN, HU, KR, SG, SI, US
         RW: BE, CH, DE, ES, FR, GB, IE, IT, NL
                            19971125
                                                            19960510
     JP 09301938
                      A2
                                           JP 1996-116545
                            19971120
    CA 2254972
                      AΑ
                                           CA 1997-2254972
                                                            19970508
                            19990324
                                                            19970508
                      A1
                                           EP 1997-918383
     EP 903337
        R: BE, CH, DE, ES, FR, GB, IT, LI, NL, IE
                            19990602
                                           CN 1997-194529
                                                            19970508
     CN 1218454
                     Α
                       Α
                            20000225
                                           KR 1998-708981
                                                            19981106
     KR 2000010840
     US 6118010
                       Α
                            20000912
                                           US 1998-147255
                                                            19981110
PRIORITY APPLN. INFO.:
                                           JP 1996-116545
                                                            19960510
                                           WO 1997-JP1543
                                                            19970508
                         CASREACT 128:35025
OTHER SOURCE(S):
    A simple, efficient and high-productivity process for prepg.
     high-quality N-(1-alkoxycarbonyl-3-phenylpropyl)amino acid derivs.
     of formula PhCH2CH2CH(CO2R)-X-Y (R = alkyl; X = Ala, Gly, Leu, Ile,
     Val, Orn or Lys or Hly with .omega.-amino group being protected with
     acyl-type protecting group; Y = OH, Ala, Gly, Leu, Ile, Val, Pro, Q,
     Q1, Q2, etc.) contg. few impurities comprises catalytic redn. of
     N-(1-alkoxycarbonyl-3-oxo-3-phenylpropyl)amino acid derivs. of
     formula PhCOCH2CH(CO2R)-X-Y (R, X, Y = same as above) in the
     presence of .gtoreq.3 equiv strong acid per 1 equiv of the
     1-alkoxycarbonyl-3-oxo-3-phenylpropyl deriv. in an alc. or a solvent
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contg. an alc. with the strong acid concn. of 0.4-5N. This process
suppresses the formation of side products, i.e. 1-alkoxycarbonyl-3-
cyclohexylpropyl derivs. Thus, 10.0 g N-[(1RS)-1-ethoxycarbonyl-3-
oxo-3-phenylpropyl]-L-alanine was added to an aq. EtOH (105 mL)
contg. 1.9N H2SO4 and 7 wt.% H2O, followed by adding 5% Pd-C contg.
50 wt.% H2O, and the resulting mixt. was hydrogenated under normal
pressure H at 20.degree. and stirring intensity 0.5-1 KW/m3. When H
was absorbed at >90% of the required quantity, the feeding of H was
stopped and the reaction mixt. was worked up to give 75%
N-[(1S)-1-ethoxycarbonyl-3-phenylpropyl]-L-alanine of 99.3 wt.%
purity contg. 0.1 wt.% N-(1-ethoxycarbonyl-3-cyclohexylpropyl)-L-
alanine and <0.1 wt.% N-(1-carboxy-3-phenylpropyl)-L-alanine.
ICM C07C229-36
ICS C07C227-16; C07K005-062; C07K005-068; C07K005-078; C07K001-113
34-3 (Amino Acids, Peptides, and Proteins)
alkoxycarbonylphenylpropyl amino acid prepn;
alkoxycarbonyloxophenylpropyl amino acid catalytic hydrogenation
Hydrogenation catalysts
   (palladium; prepn. of N-(1-alkoxycarbonyl-3-phenylpropyl)amino
   acid derivs. by catalytic hydrogenation of N-(1-alkoxycarbonyl-3-
   oxo-3-phenylpropyl)amino acid derivs.)
Hydrogenation
   (prepn. of N-(1-alkoxycarbonyl-3-phenylpropyl)amino acid derivs.
   by catalytic hydrogenation of N-(1-alkoxycarbonyl-3-oxo-3-
   phenylpropyl)amino acid derivs.)
84324-12-9P, N-[(R)-1-Ethoxycarbonyl-3-phenylpropyl]-L-alanine
199001-95-1P, N-(1-Ethoxycarbonyl-3-cyclohexylpropyl)-L-alanine
199001-96-2P, N-(1-Carboxy-3-phenylpropyl)-L-alanine
                                                       199001-97-3P,
N2-(1-Carboxy-3-phenylpropyl)-N6-trifluoroacetyl-L-lysine
199001-99-5P, N2-(1-Ethoxycarbonyl-3-cyclohexylpropyl)-N6-
                          199002-00-1P, N2-(1-Ethoxycarbonyl-3-
trifluoroacetyl-L-lysine
cyclohexylpropyl)-N6-trifluoroacetyl-L-lysyl-L-proline
199002-01-2P, N2-(1-Carboxy-3-phenylpropyl)-N6-trifluoroacetyl-L-
lysyl-L-proline
RL: BYP (Byproduct); PREP (Preparation)
   (prepn. of N-(1-alkoxycarbonyl-3-phenylpropyl)amino acid derivs.
   by catalytic hydrogenation of N-(1-alkoxycarbonyl-3-oxo-3-
   phenylpropyl)amino acid derivs.)
7440-05-3, Palladium, uses
RL: CAT (Catalyst use); USES (Uses)
   (prepn. of N-(1-alkoxycarbonyl-3-phenylpropyl) amino acid derivs.
   by catalytic hydrogenation of N-(1-alkoxycarbonyl-3-oxo-3-
   phenylpropyl) amino acid derivs.)
10009-20-8, N.omega.-Trifluoroacetyl-L-lysine
                                                15121-89-8, Ethyl
                               199002-03-4, N-(1-Ethoxycarbonyl-3-
trans-.beta.-benzoylacrylate
                                199002-04-5
oxo-3-phenylpropyl)-L-alanine
RL: RCT (Reactant); RACT (Reactant or reagent)
   (prepn. of N-(1-alkoxycarbonyl-3-phenylpropyl)amino acid derivs.
   by catalytic hydrogenation of N-(1-alkoxycarbonyl-3-oxo-3-
   phenylpropyl) amino acid derivs.)
199002-02-3P, N2-(1-Ethoxycarbonyl-3-oxo-3-phenylpropyl)-N6-
                           199611-78-4P, N2-(1-Ethoxycarbonyl-3-
trifluoroacetyl-L-lysine
phenylpropyl)-N6-trifluoroacetyl-L-lysyl-L-proline
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
RACT (Reactant or reagent)
   (prepn. of N-(1-alkoxycarbonyl-3-phenylpropyl)amino acid derivs.
   by catalytic hydrogenation of N-(1-alkoxycarbonyl-3-oxo-3-
   phenylpropyl)amino acid derivs.)
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76547-98-3P, N2-[(S)-1-Carboxy-3-phenylpropyl]-L-lysyl-L-proline
             199001-98-4P, N2-(1-Ethoxycarbonyl-3-phenylpropyl)-N6-
82717-96-2P
trifluoroacetyl-L-lysine
```

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of N-(1-alkoxycarbonyl-3-phenylpropyl)amino acid derivs. by catalytic hydrogenation of N-(1-alkoxycarbonyl-3-oxo-3phenylpropyl)amino acid derivs.)

L18 ANSWER 9 OF 17 MARPAT COPYRIGHT 2003 ACS

ACCESSION NUMBER:

126:31356 MARPAT

TITLE:

Preparation of carboxylic .alpha.-N-sulfino cyclic anhydrides as ACE inhibitor intermediates

INVENTOR(S):

Serra Mortes, Sonia; Palomo Coll, Alberto; Zupet, Rok

PATENT ASSIGNEE(S):

Centro Genesis Para La Investigacion, S.L., Spain; Serra Mortes, Sonia; Palomo Coll,

Alberto; Zupet, Rok

SOURCE:

PCT Int. Appl., 21 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.	KIND	DATE APPLICATION NO. DATE									
WO 9633					WO 1996-SI9							
W:	AL, AM,	AT, AU,	, AZ, BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,
•			, GE, HU,									
	LS, LT,	LU, LV	, MD, MG,	MK,	MN,	MW,	MX,	NO,	NΖ,	PL,	PT,	RO,
		SE, SG										
RW:			, SZ, UG,									
	GR, IE,	IT, LU	, MC, NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,
	GN											*
CA 2203	435	AA	19961031		C	A 19	96-2	2034	35	1996	0422	
AU 9652	944	A1	19961118		A	U 19	96-5	2944		1996	0422	
PRIORITY APP	LN. INFO	. :			S	I 19	95-1	40		1995	0424	
					M	0 19	96-S	Ι9		1996	0422	
OTHER SOURCE	(S):	CAS	SREACT 12	6:31	356							

AB Title compds. [I; R1 = CH2CH2Ph, or Pr; R2 = Me or (CH2)3NHR3; R3 = amino-protective group] were prepd. as intermediates in the synthesis of ACE inhibitors, esp. of enalapril and trandolapril. Thus, imidazole was chlorosulfinated with SOC12 and the product used in situ to sulfinate N-[(S)-1-ethoxycarbonyl-3-phenylpropyl]-Lalanine to give I (R1 = CH2CH2Ph, R2 = Me). The latter was amidated in situ by silylated L-proline to give enalapril.

> 308-4994 Shears Searcher :

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IÇ
     ICM C07D291-04
     ICS A61K031-41; C07D207-16; C07D209-42; A61K031-40; C07K005-06
CC
     28-10 (Heterocyclic Compounds (More Than One Hetero Atom))
     Section cross-reference(s): 34
ST
     carboxylic sulfino anhydride ACE inhibitor intermediate
     184346-33-6P
TT
     RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic
     preparation); PREP (Preparation); RACT (Reactant or reagent)
        (prepn. of carboxylic .alpha.-N-sulfino cyclic anhydrides as ACE
        inhibitor intermediates)
ΙT
     76095-16-4P, Enalapril maleate
                                      108449-50-9P
                                                     184346-34-7P
     184489-54-1P
                    184489-55-2P 184489-56-3P
                                                 184489-57-4P
     RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
     (Preparation)
        (prepn. of carboxylic .alpha.-N-sulfino cyclic anhydrides as ACE
        inhibitor intermediates)
IT
                                      80875-98-5
                                                   82717-96-2
     147-85-3, L-Proline, reactions
     145438-94-4
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of carboxylic .alpha.-N-sulfino cyclic anhydrides as ACE
        inhibitor intermediates)
L18 ANSWER 10 OF 17
                      MARPAT COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                         125:34172 MARPAT
TITLE:
                         Preparation of angiotensin converting enzyme
                         inhibitors.
INVENTOR(S):
                         Serra Mortes, Sonia; Palomo Coll, Alberto;
                         Zupet, Rok
PATENT ASSIGNEE(S):
                         Lek, Tovarna Farmacevtskih in Kemicnih Izedlkov,
                         D. D., Slovenia
SOURCE:
                         PCT Int. Appl., 47 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
    PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
                                                            DATE
                                           _____
                      ____
                           _____
                                                            -----
                            19960201
                                           WO 1995-SI17
    WO 9602564
                                                            19950713
                      A1
        W: AM, AU, BB, BG, BR, BY, CA, CN, CZ, EE, FI, GE, HU, IS, JP,
             KG, KP, KR, KZ, LK, LR, LT, LV, MD, MG, MN, MX, NO, NZ, PL,
             RO, RU, SG, SK, TJ, TM, TT, UA, US, UZ, VN
         RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE,
             IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML,
             MR, NE, SN, TD, TG
    CA 2170872
                       AΑ
                            19960201
                                           CA 1995-2170872
                                                            19950713
    AU 9529424
                            19960216
                                           AU 1995-29424
                       Α1
                                                            19950713
    EP 719280
                            19960703
                                           EP 1995-925228
                       Α1
                                                            19950713
    EP 719280
                       В1
                            19991110
        R: GB
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US 5789597

PRIORITY APPLN. INFO.:

US 1996-596214

SI 1994-290

SI 1994-450

WO 1995-SI17

19960215

19940713

19941221

19950713

19980804

Α

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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
     Title compds.(I; R = Q1-Q4, etc.), were prepd. by treatment of I (R
     = OH) (II) with ClSCOR1 or R1SOR1 (R1 = heterocyclyl such as
     imidazolyl, benzimidazolyl, 2-methylimidazolyl, triazolyl) to give
     intermediates (III) or (IV), which were reacted with (preferably
     silylated) amino acids. Thus, II was added to a soln. prepd. from
     SOC12 and imidazole in CH2Cl2 and the mixt. was stirred at -15 to
     20.degree.; the mixt. was filtered and treated with a soln. prepd.
     from Me3SiCl, proline, and Et3N in CH2Cl2. Solvent was removed and
     the residue was treated with aq. NaCl, EtOAc, and 35% HCl; the sepd.
     org. phase was treated with maleic acid to give 81.4% enalapril
     maleate.
IC
     ICM C07K005-02
     ICS C07K233-61
CC
     34-3 (Amino Acids, Peptides, and Proteins)
ST
     enalapril prepn; diquinalapril prepn; angiotensin converting enzyme
     inhibitor prepn
IT
     Peptides, preparation
     RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
     (Preparation)
        (analogs, prepn. of angiotensin converting enzyme inhibitors)
TΥ
     151387-05-2P
     RL: BYP (Byproduct); PREP (Preparation)
        (prepn. of angiotensin converting enzyme inhibitors)
IT
     76095-16-4P, Enalapril maleate 138332-08-8P, Enalapril
                    149404-21-7P, Enalapril sodium 177696-02-5P
     hydrochloride
     177696-03-6P
     RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
     (Preparation)
        (prepn. of angiotensin converting enzyme inhibitors)
TΤ
     147-85-3, Proline, reactions 77497-95-1
                                               82717-96-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of angiotensin converting enzyme inhibitors)
IT
     7364-47-8P
                 82006-94-8P
                               129258-49-7P
                                             177545-70-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent).
        (prepn. of angiotensin converting enzyme inhibitors)
L18 ANSWER 11 OF 17 MARPAT COPYRIGHT 2003 ACS
                         118:154603 MARPAT
ACCESSION NUMBER:
TITLE:
                         Method of treating premenstrual syndrome by
                         administration of an angiotensin-converting
                         enzyme inhibitor
INVENTOR(S):
                         DePadova, Anthony Salvator
PATENT ASSIGNEE(S):
                         Warner-Lambert Co., USA
SOURCE:
                         PCT Int. Appl., 18 pp.
                         CODEN: PIXXD?
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND
                            DATE
                                           APPLICATION NO.
                                                            DATE
     WO 9302679
                       Α1
                            19930218
                                           WO 1992-US6210
                                                            19920803
```

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W: AU, CA, JP
        RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, SE
                     A1 19930302
                                          AU 1992-24139
                                                           19920803
    AU 9224139
                                          US 1991-740557
                                                           19910805
PRIORITY APPLN. INFO.:
                                          WO 1992-US6210
                                                           19920803
    A method of treating premenstrual syndrome is described which
AB
    comprises the administration of a daily dose of an effective amt. of
    an angiotensin-converting enzyme inhibitor, e.g. quinapril, to a
    female of menstrual age. Tablet formulations of quinapril-HCl are
    included.
    ICM A61K031-475
TC
    ICS A61K031-40; A61K031-675
CC
    63-6 (Pharmaceuticals)
    premenstrual syndrome angiotensin converting enzyme inhibitor;
ST
    tablet quinapril hydrochloride premenstrual syndrome
ΙT
    Ovarian cycle
        (disorder, premenstrual syndrome, treatment of, pharmaceutical
       compn. contg. angiotensin-converting enzyme inhibitor for)
    Pharmaceutical dosage forms
ΙT
        (tablets, of quinapril hydrochloride, for premenstrual syndrome
       treatment)
    9015-82-1, Angiotensin-converting enzyme
TΤ
    RL: BIOL (Biological study)
        (inhibitors of, pharmaceutical compn. contg., for premenstrual
       syndrome treatment)
                                                    75847-73-3,
IT
     62571-86-2, Captopril
                            74258-86-9, Alacepril
                76547-98-3, Lisinopril 80876-01-3, Indolapril
                             82586-55-8, Quinapril hydrochloride
    81872-10-8, Zofenopril
                             82924-03-6, Pentopril 83435-66-9,
    82834-16-0, Perindopril
               83647-97-6, Spirapril 85441-61-8, Quinapril
                 87333-19-5, Ramipril 88768-40-5, Cilazapril
    86541-75-5
                             109214-55-3, Libenzapril
    98048-97-6, Fosinopril
    RL: BIOL (Biological study)
        (pharmaceutical compn. contg., for premenstrual syndrome
       treatment)
L18 ANSWER 12 OF 17
                     MARPAT COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                        117:172104 MARPAT
                        Methods for the synthesis of aminosuberic acid
TITLE:
                        derivatives
                        Hoffmann, Gerhard; Liedtke, Bernhard; Vollmer,
INVENTOR(S):
                        Karl Otto
                        Goedecke AG, Germany
PATENT ASSIGNEE(S):
                        Ger. Offen., 6 pp.
SOURCE:
                        CODEN: GWXXBX
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                        German
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                     KIND DATE
                                                           DATE
                                          APPLICATION NO.
    PATENT NO.
                                          ______
    _____
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                           _____
    ,DE 4037960
                           19920604
                                          DE 1990-4037960 19901129
                      Α1
                                          DE 1990-4037960 19901129
PRIORITY APPLN. INFO.:
                        CASREACT 117:172104
OTHER SOURCE(S):
    For diagram(s), see printed CA Issue.
GΙ
    Aminosuberic acid derivs. I (R1 = alkyl or alkenyl with up to 6
AB
    carbon atoms, C5-7-cycloalkyl, C5-7-cycloalkenyl,
```

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C7-12-cycloalkylalkyl, C6-10-aryl, C7-14-aralkyl, mono- or bicyclic
    heterocyclic group; R2 = aryl; R3 = C1-6-alkyl, C2-6-alkenyl,
    C7-14-aralkyl; Z forms a heterocyclic ring) were prepd. by the
     selective cleavage of PhCH2O2CCHR1NHCH(CO2R3)CH2COR2 with AlCl3,
     condensing the resulting HO2CCHR1NHCH(CO2R3)CH2COR2 with
    heterocyclic compd. II, and hydrogenating the resulting ketone III
    with H2, deuterium or tritium. Thus, (S,S)-PhCOCH2CH(CO2Et)-Ala-
    OCH2Ph was debenzylated with AlCl3 in the presence of anisole in
    CH2Cl2/MeNO2 to give 86% (S,S)-PhCOCH2CH(CO2Et)-Ala-OH, which was
    condensed with (S)-1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid
    benzyl ester by DCC/1-hydroxybenzotriazole in CH2Cl2 to give product
    IV (isolated as the HCl salt). IV was hydrogenated over Pd/C in the
    presence of HCl to give 88% tetrahydroisoquinoline V.HCl.
    ICM C07H005-06
     ICS C07D217-26
    34-2 (Amino Acids, Peptides, and Proteins)
     Section cross-reference(s): 1
     aminosuberic acid; suberic acid amino
     77497-96-2
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (coupling of, with alanine deriv.)
     87269-98-5
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (debenzylation of)
     143442-61-9P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and conversion of, to hydrochloride salt)
     87269-99-6P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
    RACT (Reactant or reagent)
        (prepn. and coupling of, with tetrahydroisoquinolinecarboxylic
        acid benzyl ester)
     143381-50-4P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
    RACT (Reactant or reagent)
        (prepn. and hydrogenation or deuteration of)
    82586-55-8P
                   143381-51-5P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
    RACT (Reactant or reagent)
        (prepn. and sapon. of)
     3054-07-7DP, 2-Aminosuberic acid, derivs.
     143381-52-6P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
L18 ANSWER 13 OF 17
                      MARPAT COPYRIGHT 2003 ACS
                         117:143453 MARPAT
ACCESSION NUMBER:
                         Use of a combination of an ACE
TITLE:
                         (angiotensin-converting enzyme) inhibitor with a
                         calcium antagonist in the treatment of
                         proteinuria
                         Becker, Reinhard; Henning, Rainer; Teetz,
INVENTOR(S):
                         Volker; Urbach, Hansjoerg
PATENT ASSIGNEE(S):
                         Hoechst A.-G., Germany
SOURCE:
                         Eur. Pat. Appl., 22 pp.
                         CODEN: EPXXDW
DOCUMENT TYPE:
                         Patent .
LANGUAGE:
                         German
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FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

				KIND	DATE		APP	LICATION NO.	DATE
	EP EP	488059		A3	19921125		EP	1991-119892	19911121
	ĒΡ	488059		B1	19950906	ED	CD C	וון דו חדמי	MI SE
		R: AT,	BE,	CH, DE,	DK, ES,	rK,	GD, G	R, IT, LI, LU,	10011121
	EP	649654		AI	19950426		гF	1994-117179	19911121
	EP	649654		BI	19990210	DD.	CD C	וז דו חד מי	NI CE
		R: AT,	BE,	CH, DE,	DK, ES,	rk,	GB, G	R, IT, LI, LU,	10011121
	EŞ	2079545		T3	19960116		ES.	1991-119892	19911121
	ΑT	176592 2129563		E	19990215		AT	1994-117179	19911121
	ES	2129563		Т3	19990616		ES	1994-117179 1991-88117	19911121
	ΑU	9188117		Al	19920528		AU	1991-88117	19911126
	ΑU	655784		B2	19950112				
	ÇA	2055948		AA	19920528		CA		19911126
	NO	9104637		A	19920529		иО	1991-4637	19911126
	ZA	9109318		Α	19920826		ZA	1991-9318	19911126
	JΡ	04308533		A2 ·	19921030			1991-310608	
	HU	62468		A2	19921030 19930528 20010428		HU	1991-3674	19911126
	HU	219447		В	20010428				
	CN	1072601		A	19930602		CN	1991-111099	19911126
		1060679		В	20010117			•	
	US	5236933		Ā	19930817		US	1991-798501	19911126
	SK	279626			19990111		SK	1991-3587	19911126
	CZ	286168		В6	20000216		CZ	1991-3587	19911126
	US	5366994		Α	19941122		US	1993-57516	19930506
	CZ.	286187		В6	19941122 20000216		CZ	1997-2830	19970908
	HK	1011927		Ä1	20000728			1998-98113023	19981209
PRIOR		APPLN.					DE	1990-4037691	19901127
11(101				-			EP	1991-119892	19911121
	•						CS	1991-3587	19911126
							US	1991-798501	19911126
מ ת	71	ACE inhi	hitor	P302C	CHRANRSC (·0)C		CH (CO2R2) (CH2)	

An ACE inhibitor R3O2CCHR4NR5C(:0)CHR1NHCH(CO2R2)(CH2)nR [n = 1, 2; AB $R = H_{\lambda}$ (substituted) aliph., alicyclic, arom., hydrocarbyl- or heterocyclyloxy or -thio; R1 = H, (substituted) hydrocarbyl or heteroarom.; R2, R3 = H, (substituted) aliph., alicyclic, arom., araliph.; R4 and R5 complete a heterocyclic mono-, bi-, or tricyclic ring system with 3-15 C atoms], combined with a Ca antagonist, is used for prevention and therapy of proteinuria secondary to diabetes mellitus, glomerulosclerosis, and loss of kidney mass. Thus, rats with 1 kidney removed and the other infarcted through ligation were administered ramipril (ACE inhibitor; 1.4 mg/kg) and felodipine (Ca antagonist; 41 mg/kg) in the feed. An increase in proteinuria from <20 to 105 mg/24 h was obsd. in controls, compared to only 31 mg/24 h in treated rats. Tablets were prepd. contg. trandolapril (ACE inhibitor) 3, verapamil (Ca antagonist) 50, corn starch 130, gelatin 8.0, microcryst. cellulose 2.0, and Mg stearate 2.0 g/1000.

IC ICM A61K037-64 ICS A61K045-06

ICI A61K037-64, A61K031-135; A61K037-64, A61K031-44; A61K037-64, A61K031-55

CC 1-8 (Pharmacology)

Section cross-reference(s): 63

ST angiotensin converting enzyme inhibitor proteinuria; calcium antagonist proteinuria treatment

```
ΙT
     Proteins, biological studies
     RL: BIOL (Biological study)
        (metabolic disorders, proteinuria, treatment of, with
       angiotensin-converting enzyme inhibitor and calcium antagonist)
IT
     7440-70-2, Calcium, biological studies
     RL: BIOL (Biological study)
        (antagonist of, proteinuria treatment with angiotensin-converting
       enzyme inhibitor and)
ΙT
     9015-82-1, Angiotensin-converting enzyme
     RL: BIOL (Biological study)
        (inhibitor of, proteinuria treatment with calcium antagonist and)
IT
     119884-28-5
                  119920-75-1 119920-79-5 143375-98-8 143375-99-9
     143376-00-5
                  143376-01-6
     RL: BIOL (Biological study)
        (proteinuria treatment with)
     52-53-9, Verapamil 21829-25-4, Nifedipine
IT
                                                 42399-41-7
     RL: BIOL (Biological study)
       (proteinuria treatment with angiotensin-converting enzyme
       inhibitor and)
IT
     91-21-4D, derivs.
                        123-75-1D, Pyrrolidine, derivs.
                                                         172-62-3D,
              175-94-0D, 2-Azaspiro[4.4]nonane, derivs.
                                                         176-66-9D,
     2-Azaspiro[4.5]decane, derivs. 504-78-9D, Thiazolidine, derivs.
     4375-14-8D, Octahydroindole, derivs. 5661-02-9D, derivs.
     5661-03-0D, derivs. 6329-61-9D, Decahydroisoquinoline, derivs.
     7140-62-7D, Decahydrocyclohepta[b]pyrrole, derivs.
                                                        21850-12-4D,
              27202-71-7D, 2-Azabicyclo[3.1.0]hexane, derivs.
     62571-86-2, Captopril 73263-16-8D, Spiro[(bicyclo[2.2.2]octane)-
     2,3'-pyrrolidine], derivs. 74258-86-9, Alacepril 75847-73-3,
                76547-98-3, Lisinopril 81872-10-8, Zofenopril
     Enalapril
                83435-66-9, Delapril 83647-97-6, Spirapril
     82834-16-0
    84768-09-2 85441-61-8, Quinapril 85856-54-8, Moveltipril
    86541-75-5, Benazepril 87333-19-5, Ramipril 87679-37-6,
                                89371-37-9
                  88768-40-5
                                           90130-77-1, S 9650
     Trandolapril
     98048-97-6, Fosinopril 103775-10-6, Moexipril 109214-55-3,
     Libenzapril 109683-61-6, FPL 63547 110221-44-8 111223-26-8,
    Ceranapril 143384-22-9D, derivs. 143442-70-0D, derivs.
     144275-78-5D, derivs.
    RL: BIOL (Biological study)
        (proteinuria treatment with calcium antagonist and)
L18 ANSWER 14 OF 17 MARPAT COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                        114:88663 MARPAT
                        Combinations of angiotensin-converting enzyme
TITLE:
                        (ACE) inhibitor and carbonic anhydrase (CA)
                        inhibitor for treating glaucoma
                        Lotti, Victor J.; Baldwin, John J.
INVENTOR(S):
PATENT ASSIGNEE(S):
                        Merck and Co., Inc., USA
SOURCE:
                        Eur. Pat. Appl., 17 pp.
                        CODEN: EPXXDW
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO.
                     KIND
                           DATE
                                         APPLICATION NO.
                                          _____
     ______
                           -----
                                     EP 1989-313161
                           19900627
    EP 375299 A1
                                                          19891215
        R: AT, BE, CH, DE, ES, FR, GB, IT, LI, LU, NL, SE
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CA 1989-2005754 19891218 CA 2005754 19900619 AA19900620 DK 1989-6412 19891218 DK 8906412 Α A2 19900823 JP 1989-329369 19891219 JP 02212425 US 1988-285932 19881219 PRIORITY APPLN. INFO.:

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

```
The ACE inhibitors are carboxylates RCHR5COACO2R6 [A = Q, Q1, Q2, Q3; R = R2CH2CR4(CO2R3)NH, HSCH2; R1, R3, R4, R6 = H, alkyl; R2 = alkyl, PhCH2, PhO, PhS, PhCH2O; R5 = R1, aminoalkyl; p, q = 0-2; r = 1, 2]. The CA inhibitors are sulfonamides I (R7 = H, R8 = Q4; or R7R8 = Q5, Q6, etc.; R9, R10 = H, alkyl; R11 = H, alkanoyl; R12 = alkoxyalkyl; R13 = H, alkoxyalkoxyalkyl). Compns. comprising an ACE inhibitor and a CA inhibitor are synergistic ophthalmic drugs for the treatment of glaucoma (no data).
```

IC ICM A61K037-64 ICS A61K009-06

ICI A61K037-64, A61K031-38

CC 63-6 (Pharmaceuticals)

ST glaucoma synergism drug enzyme inhibitor; angiotensin converting enzyme inhibitor glaucoma; carbonic anhydrase inhibitor glaucoma

IT Glaucoma (disease)

(treatment of, synergistic drugs contg. angiotensin-converting enzyme inhibitors and carbonic anhydrase inhibitors for)

IT 63250-36-2D, mixts. with carbonic anhydrase inhibitors 76391-23-6D, mixts. with carbonic anhydrase inhibitors 83601-86-9D, mixts. with carbonic anhydrase inhibitors 119731-42-9D, mixts. with angiotensin-converting enzyme inhibitors 122266-90-4D, mixts. with angiotensin-converting enzyme inhibitors 128620-92-8D, mixts. with angiotensin-converting enzyme inhibitors

131898-30-1D, mixts. with carbonic anhydrase inhibitors

RL: BIOL (Biological study)

(for glaucoma treatment, synergistic)

IT 9001-03-0, Carbonic anhydrase

RL: BSU (Biological study, unclassified); BIOL (Biological study) (inhibitors, mixts. with angiotensin-converting enzyme inhibitors, synergistic, for glaucoma treatment)

IT 9015-82-1, Angiotensin-converting enzyme

RL: BSU (Biological study, unclassified); BIOL (Biological study) (inhibitors, mixts. with carbonic anhydrase inhibitors, synergistic, for glaucoma treatment)

L18 ANSWER 15 OF 17 MARPAT COPYRIGHT 2003 ACS

ACCESSION NUMBER:

111:120883 MARPAT

TITLE:

Antihypertensive pharmaceuticals containing 1,4-dihydropyridine-3,5-dicarboxylic acid

analogs and isoquinoline carboxylic acid analogs

INVENTOR(S): Strosberg, Arthur M.; Whiting, Roger L.

PATENT ASSIGNEE(S):

Syntex (U.S.A.), Inc., USA

SOURCE:

Eur. Pat. Appl., 13 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	TENT 1	NO.		KIN	1D	DATE			AP	PLIC	CATI	ON N	ο.	DATE	
	EP	25983	 38		A2	2	1988	0316		EP	198	 37-1	 1312	1	19870	908
	ΕP	25983	38		A3	3	1989	1115								
		'R:	AT,	BE,	CH,	DE,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE	•
	DK	87046	690		Α		1988	0310				37-4			19870	908
	ΑU	87783	157		A1	L	1988	0317		ΑU	198	37-7	8157		19870	908
	JΡ	63079	9831		A2	2	1988	0409		JP	198	37-2	2514	1	19870	908
	ZA	8706	716		Α		1989	0426		z_A	198	37-6	716		19870	908
PRIOR	RITY	APPI	LN.	INFO.	:					US	198	36-9	0536	1	19860	909
GT																

Ι

Pharmaceuticals contain 2,6-dimethyl-4-phenyldihydropyridine-3,5-AB dicarboxylic acid derivs. (I; R1 = o- or m-substituted NO2, CF3, halo; R2 = alkyl, CH2CH2OMe; A = alkylene; R3 = alkyl, alkoxy, or optionally substituted Ph, or phenylalkyl; R4, R5 = H, alkyl) or salts of I, and an isoquinolinecarboxylic acid deriv. (II; R = H, alkyl) or salts of II. Antihypertensive tablets contained 2,6-dimethyl-4-(3-nitrophenyl)-1,4-dihydropyridine-3,5-dicarboxylic acid-3-Me ester-5-.beta.-(N-benzyl-N-methylamino)-Et ester (nicardipine) (III) (10 mg) and 2-[2-[(1-methoxycarbonyl)-3phenylpropyl]amino]-1-oxopropyl-1,2,3,4-tetrahydro-6,7-dimethoxy-3isoquinolinecarboxylic acid (IV) (20 mg), corn starch (20 mg) spray-dried lactose (153 mg), and Mg stearate (2 mg). A combination of III (3 mg/kg orally) and IV (10 mg/kg orally) induced a greater antihypertensive effect on renal hypertensive rats with a systolic blood pressure >160 mmHg than did III or IV alone; the potentiation effect lasted >4 h.

IC ICM A61K031-47

```
ICI A61K031-47, A61K031-44
     63-6 (Pharmaceuticals)
CC
     Section cross-reference(s): 1
     pyridine isoquinolinecarboxylate antihypertensive; calcium channel
ST
     blocker antihypertensive
     Antihypertensives
ΙT
        (calcium channel blocker-angiotensin-converting enzyme inhibitor
        mixts. as)
IT
     Ion channel blockers
        (calcium, mixts. with angiotensin-converting enzyme inhibitors,
        as antihypertensive agents)
ΤT
     122441-13-8
     RL: BIOL (Biological study)
        (antihypertensive agent)
     55985-32-5D, mixts. with isoquinolinecarboxylate analogs
IT
     122379-46-8D, mixts. with dihydropyridine analogs
     RL: BIOL (Biological study)
        (antihypertensive agents)
IT
     9015-82-1, Angiotensin-converting enzyme
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitors, mixts. with calcium channel blockers, as
        antihypertensive agents)
L18 ANSWER 16 OF 17 MARPAT COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                         110:141552 MARPAT
                         Pharmaceuticals containing angiotensin-
TITLE:
                         converting enzyme inhibitors and ascorbates as
                         stabilizers
                         Murthy, Kuchi Sury; Harris, Michael Ray;
INVENTOR(S):
                         Hokanson, Gerard Clifford; Reisch, Robert
                         George, Jr.; Fawzi, Mahdi Bakir; Waldman, Frank
                         Stanley
PATENT ASSIGNEE(S):
                         Warner-Lambert Co., USA
SOURCE:
                         Eur. Pat. Appl., 6 pp.
                         CODEN: EPXXDW
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
                         1
PATENT INFORMATION:
```

PATENT I	NO. KINI	DATE	API	PLICATION NO.	DATE
EP 2648	87 A1 87 B1		EP	1987-115281	19871019
R:	AT, BE, CH, I	DE, ES, FR,	GB, GR, I	IT, LI, LU, N	IL, SE
US 4830	853 A	19890516	US	1986-921717	19861020
CA 1297	024 A1	19920310	CA	1987-547231	19870918
ZA 8707	132 A	19890426	ZA	1987-7132	19870922
AU 8779	397 A1	19880421	AU	1987-79397	19871006
AU 6040	61 B2	19901206			
DK 8705	435 A	19880421	DK	1987-5435	19871016
FI 8704	594 A	19880421	FI	1987-4594	19871019
NO 8704	352 A	19880421	NO	1987-4352	19871019
JP 6310	4931 A2	19880510	JP	1987-261931	19871019
JP 2703	906 B2	19980126			
AT 6709	0 E	19910915	AT	1987-115281	19871019
ES 2040	726 T3	19931101	ES	1987-115281	19871019
PRIORITY APP	LN. INFO.:		US	1986-921717	19861020

EP 1987-115281 19871019 AΒ Pharmaceuticals contain an angiotensin converting-enzyme (ACE) inhibitor which is susceptible to discoloration, ascorbic acid (I) and/or Na ascorbate as color stabilizing agent(s), and optionally other additives. A formulation contained quinapril 5, I 20, lactose 71, and hydrogenated vegetable oil 4% by wt. This compn. was stable at 80% relative humidity for .gtoreq.24 h. ICICM A61K031-44 ICS A61K031-40; A61K047-00 CC 63-6 (Pharmaceuticals) angiotensin converting enzyme inhibitor ascorbate; antihypertensive ST ascorbate Discoloration prevention IT(of angiotensin-converting enzyme inhibitor-contg. pharmaceuticals, ascorbates for) 9015-82-1, Angiotensin converting enzyme ΙT RL: BSU (Biological study, unclassified); BIOL (Biological study) (inhibitors, pharmaceuticals contg. ascorbates and) 50-81-7, Ascorbic acid, biological studies 134-03-2, Sodium ΤТ ascorbate RL: BIOL (Biological study) (pharmaceuticals contg. antihypertensive angiotensin-converting enzyme inhibitors and) ΙT 75847-73-3, Enalapril 82768-84-1 85441-61-8, Quinapril 103775-10-6 RL: BIOL (Biological study) (pharmaceuticals contg. ascorbates and) L18 ANSWER 17 OF 17 MARPAT COPYRIGHT 2003 ACS ACCESSION NUMBER: 109:48448 MARPAT Neutral metalloendopeptidase inhibitors in the TITLE: treatment of hypertension, compositions and kits containing the inhibitors, manufacture of the compositions, compounds of the compositions and their preparation Haslanger, Martin F.; Sybertz, Edmund, Jr.; INVENTOR(S): Neustadt, Bernard R.; Smith, Elizabeth M. Schering Corp., USA PATENT ASSIGNEE(S): SOURCE: Eur. Pat. Appl., 167 pp. CODEN: EPXXDW DOCUMENT TYPE: Patent LANGUAGE: · English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE APPLICATION NO. DATE PATENT NO. EP 1987-108730 19870617 EP 254032 A2 19880127 ΑЗ 19900905 EP 254032 R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE US 4749688 Α 19880607 US 1986-876610 19860620 US 1987-32153 US 4801609 Α 19890131 19870327 19931020 EP 1993-107499 EP 566157 A1 19870617 R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE FI 8702720 Α 19871221 FI 1987-2720 19870618

Searcher: Shears 308-4994

AU 8774458

AU 602701

ZA 8704413

Α1

В2

19871224

19901025

19880224

AU 1987-74458

ZA 1987-4413

19870618

19870618

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HU 1987-2786
                                                             19870618
     HU 44940
                       Α2
                            19880530
     IL 82908
                       A1
                            19910916
                                           IL 1987-82908
                                                             19870618
                       Α
                                           DK 1987-3138
                                                             19870619
     DK 8703138
                            19871221
                       Α
                            19871221
                                           NO 1987-2589
                                                             19870619
     NO 8702589
                       A2
                            19880220
                                           JP 1987-153219
                                                             19870619
     JP 63039855
                       B2
                            19961009
     JP 2542620
     JP 08283153
                       A2
                            19961029
                                           JP 1995-246555
                                                             19870619
     US 5061710
                       Α
                            19911029
                                           US 1987-133669
                                                             19871216
                                           AU 1990-68517
    AU 9068517
                       Α1
                            19910718
                                                             19901227
    AU 636423
                       B2
                            19930429
                                           US 1991-90002282 19910214
     US 4801609
                       В1
                            19931109
                                           US 1991-741025
     US 5262436
                       Α
                            19931116
                                                             19910806
     JP 08176100
                       A2
                            19960709
                                           JP 1995-246554
                                                             19950821
PRIORITY APPLN. INFO.:
                                           US 1986-876610
                                                             19860620
                                           US 1987-32153
                                                             19870327
                                                             19870617
                                           EP 1987-108730
                                           JP 1987-153219
                                                             19870619
                                           US 1987-133669
                                                             19871216
AB
    Neutral metalloendopeptidase (NMEP) inhibitor is used alone or
     combined with an atrial peptide or an angiotensin converting enzyme
     (ACE) inhibitor for prepn. of pharmaceutical compns. for treating
     hypertension. The compns. are obtained by mixing a NMEP inhibitor,
     alone or combined with an atrial peptide or ACE inhibitor, with a
     pharmaceutically acceptable carrier. S-(4-Methylbenzyl)-L-cysteine,
    Me ester hydrochloride was prepd. by adding thionyl chloride
     dropwise to N-tert-butyloxycarbonyl-S-(4-methylbenzyl)-L-cysteine in
    MeOH, heating the mixt. under reflux for 90 min, cooling to room
     temp., and concg. in vacuo. Rats with induced hypertension were
     dosed s.c. with N-(N-[L-1-(2,2-dimethyl-1-
     oxopropoxy)methoxy]carbonyl)-2-phenylethyl)-L-phenylalanine]-.beta.-
     alanine and 1-[(2S)-3-mercapto-2-methyl-1-oxypropyl]-L-proline in Me
    cellulose vehicle to give a 1-, 2-, 3-, and 4-h decrease in blood
    pressure of 14, 19, 19, and 15 mMHg vs. an increase of 14, 11, 11,
     and 8 with the NMEP inhibitor alone and a decrease of 11, 7, 1, and
     1 mMHg with the ACE inhibitor alone.
     ICM A61K037-64
IC
CC
     1-8 (Pharmacology)
     Section cross-reference(s): 34, 63
ST
    NMEP inhibitor atrial peptide antihypertensive
IT
    Antihypertensives
        (neutral metalloendopeptidase inhibitors and angiotensin
        converting enzyme inhibitors or atriopeptins as)
ΙT
                             75847-73-3
                                          76547-98-3, Lysinopril
     62571-86-2, Captopril
                              81045-50-3, Pivalopril
                                                       81872-10-8,
     80876-01-3, Indolapril
     Zofenopril
                  82834-16-0, Perindopril
                                            82924-03-6, Pentopril
     83647-97-6, Spirapril
                             87333-19-5, Ramipril
                                                    88768-40-5,
    Cilazapril
     RL: BIOL (Biological study)
        (angiotensin converting enzyme inhibitor, antihypertensive contq.
        neutral metalloendopeptidase inhibitor and)
IT
    82586-52-5
    RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); BIOL (Biological study)
        (angiotensin converting enzyme inhibitor, antihypertensive contg.
        neutral metalloendopeptidase inhibitor and)
ΙT
     115156-91-7
                   115156-92-8
                                 115156-94-0
     RL: BIOL (Biological study)
        (angiotensin converting enzyme inhibitor-neutral
```

```
metalloendopeptidase inhibitor mixt., with antihypertensive
        properties)
TΤ
     88898-17-3
                  89139-53-7, Atriopeptin-21 (rat)
                                                      89139-54-8
     89213-87-6
                                                      90052-57-6
                  89944-37-6, Atriopeptin-33 (rat)
                  90984-99-9
                                94705-21-2, Atriopeptin 33 (human)
     90817-13-3
                  98897-20-2
                                98929-56-7
                                             98929-57-8
                                                           102686-43-1
     95896-08-5
     RL: BIOL (Biological study)
        (antihypertenive pharmaceuticals contg. neutral
        metalloendopeptidase inhibitors and)
     82707-54-8
ΙT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitors, antihypertensive compns. contg. atriopeptins or
        angiotensin converting enzyme inhibitors and)
     9015-82-1
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitors, pharmaceuticals with neutral metalloendopeptidase
        inhibitor, with antihypertensive properties)
                  95198-45-1
                                105368-09-0
                                              115156-93-9
                                                             115355-20-9
ΙT
     83861-02-3
                   115355-22-1
     115355-21-0
     RL: BIOL (Biological study)
        (neutral metalloendopeptidase inhibitor, antihypertensive contg.)
                                             95198-15-5
                                                          95198-52-0
                                95176-40-2
ΙT
     83825-63-2
                  83861-02-3
                                                115370-39-3
                                  115355-22-1
                                                               115370-40-6
     115156-93-9
                   115355-21-0
                                  115370-43-9
                                                115370-44-0
                                                               115370-45-1
     115370-41-7
                   115370-42-8
                                                115370-49-5
     115370-46-2
                   115370-47-3
                                  115370-48-4
                                                               115370-50-8
                                                115406-23-0
                                  115406-22-9
     115370-51-9
                   115370-52-0
     RL: BIOL (Biological study)
        (neutral metalloendopeptidase inhibitor, with antihypertensive
        properties)
ΙT
     85637-73-6, Atriopeptin
     RL: BIOL (Biological study)
        (pharmaceuticals with neutral metalloendopeptidase inhibitors,
        with antihypertensive properties)
                                                               115369-83-0P
ΙT
                  27650-96-0P
                                 95244-36-3P
                                               105852-67-3P
     2107-84-8P
                    115369-85-2P
                                    115369-86-3P
                                                   115369-87-4P
     115369-84-1P
                                                   115369-91-0P
     115369-88-5P
                    115369-89-6P
                                    115369-90-9P
                    115369-93-2P
                                    115369-94-3P
                                                   115369-95-4P
     115369-92-1P
     115369-96-5P
                                    115369-98-7P
                                                   115369-99-8P
                    115369-97-6P
     115370-00-8P
                                    115370-02-0P
                                                   115370-03-1P
                    115370-01-9P
                                                   115370-07-5P
     115370-04-2P
                    115370-05-3P
                                    115370-06-4P
     115370-08-6P
                    115370-09-7P
                                    115370-10-0P
                                                   115370-11-1P
                                                   115406-17-2P
     115370-53-1P
                    115370-54-2P
                                    115391-56-5P
                                                   115406-21-8P
                    115406-19-4P
                                    115406-20-7P
     115406-18-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (prepn. and reaction of, in neutral metalloendopeptidase
        inhibitor prepn.)
                                                   115370-15-5P
TT
                    115370-13-3P
                                    115370-14-4P
     115370-12-2P
     115370-16-6P
                    115370-17-7P
                                    115370-18-8P
                                                   115370-19-9P
                    115370-21-3P
                                    115370-22-4P
                                                   115370-23-5P
     115370-20-2P
                    115370-25-7P
                                    115370-26-8P
                                                   115370-27-9P
     115370-24-6P
                    115370-29-1P
                                                   115370-31-5P
     115370-28-0P
                                    115370-30-4P
     115370-32-6P
                    115370-33-7P
                                                   115370-35-9P
                                    115370-34-8P
                    115370-37-1P
     115370-36-0P
                                    115370-38-2P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of, as neutral metalloendopeptidase inhibitor)
IT
                   34805-17-9P, O-Benzyl-L-tyrosine methyl ester
     13026-12-5P
                     52844-67-4P
                                    66076-76-4P
                                                  71449-22-4P
     hydrochloride
```

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84952-45-4P 88389-36-0P
                                              88424-57-1P
                                                            88660-34-8P
    74401-75-5P
                   98574-43-7P
                                115355-23-2P
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     95260-87-0P
     115355-25-4P
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                                   115355-27-6P
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                                   115355-31-2P
                                                 115355-32-3P
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                   115355-34-5P
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     115355-33-4P
                   115355-38-9P
                                  115355-39-0P
                                                 115355-40-3P
     115355-37-8P
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                   115355-46-9P
                                  115355-47-0P
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                                                 115355-52-7P
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                                                 115405-40-8P
    115405-37-3P
                   115405-38-4P
                                  115405-39-5P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of, as neutral metalloendopeptidase inhibitor inhibitor)
                                               105-53-3
ΙT
    76-05-1, Trifluoroacetic acid, reactions
                                                         115-11-7,
     Isobutylene, reactions 121-44-8, Triethylamine, reactions
     141-52-6, Sodium ethanolate 141-82-2, Malonic acid, reactions
     507-09-5, Thioacetic acid, reactions 1535-57-5 2177-63-1
                                                3218-36-8, 4-Biphenyl
     3163-27-7, .alpha.-Bromomethylnaphthalene
     carboxaldehyde 4510-08-1, L-Methionine amide 6287-94-1
                                          13485-59-1, L-Alanyl-L-proline
     10332-17-9, Methionine methyl ester
                 42294-52-0, S-(4-Methylbenzyl)-L-cysteine 52844-67-4
     14510-12-4
                 80969-99-9 83024-49-1 84609-11-0
                                                       100484-62-6
     61925-77-7
                 115355-30-1 115369-81-8
                                              115369-82-9
     115355-23-2
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, in neutral metalloendopeptidase inhibitor prepn.)
```

FILE 'MARPATPREV' ENTERED AT 12:08:31 ON 21 APR 2003 L16 STR

REP G1=(1-3) CH2 VAR G2=OH/30 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM MLEVEL IS CLASS AT 22 26 32 GGCAT IS UNS AT 32 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RSPEC I

NUMBER OF NODES IS 32

STEREO ATTRIBUTES: NONE

ATTRIBUTES SPECIFIED AT SEARCH-TIME:

ECLEVEL IS LIM ON ALL NODES ALL RING(S) ARE ISOLATED

L19 0 SEA FILE=MARPATPREV SSS FUL L16 (MODIFIED ATTRIBUTES)

100.0% PROCESSED 3

3 ITERATIONS

0 ANSWERS

SEARCH TIME: 00.00.01

(FILE 'CASREACT' ENTERED AT 12:08:58 ON 21 APR 2003) L21 STR

REP G1=(1-3) CH2 VAR G2=OH/30 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM GGCAT IS UNS AT 32 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 32

STEREO ATTRIBUTES: NONE

L23) 4 SEA FILE=CASREACT SSS FUL L21 (32 REACTIONS)

100.0% DONE 211 VERIFIED 32 HIT RXNS 4 DOCS

SEARCH TIME: 00.00.01

L23 ANSWER 1 OF 4 CASREACT COPYRIGHT 2003 ACS

ACCESSION NUMBER:

136:279699 CASREACT

TITLE: Preparation of amino acid salts soluble in

organic solvents and their use in dipeptide

synthesis

INVENTOR(S): Palomo Nicolau, Francisco Eugenio; Palomo Coll,

Antonio Luis

PATENT ASSIGNEE(S):

Spain

SOURCE:

Span., 23 pp.

CODEN: SPXXAD

DOCUMENT TYPE:

Patent

LANGUAGE:

FAMILY ACC. NUM. COUNT:

Spanish

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ES 2156050	A1	20010601	ES 1998-753	19980331
ES 2156050	B1	20020316		
RIORITY APPLN INFO			ES 1998-753	19980331

MARPAT 136:279699 OTHER SOURCE(S):

.alpha.-Amino acids salts with org. super bases, ROCO2-A-NH-Q+-[N(B)D]n[Q=C(n=1 or 2) or P(n=3); A, B, D=alkyl,alkylaryl or may combine to form a heterocyclic group; R = H or a side chain of an amino acid; RO is an aminohetero(bi)cyclic C5-C10 residue having one or two atoms O, S or N or a group R1NHCHR, where R = H or a side chain of an optionally protected amino acid, R1 = H, Ph3C; or R and R1 may combine to form a mono-, bi-, or tricyclic heterocyclic ring], which are sol. in org. solvents, were prepd. for use in the prepn. of dipeptides. Tetramethylformamidinium, benzotriazolylmethyl, and oxime active esters were used in this process. Thus, treatment of an acetonitrile soln. of L-proline and DBU with N-[(1S)-1-(ethoxycarbonyl)-3-phenylpropyl]-L-alanine O-acetone oxime (prepn. given) afforded 95% enalapril (isolated as

$$RX(8)$$
 OF 13 $AM + AN ===> AO$

the maleate).

ΑM

ΑN

AO YIELD 90%

RX(8) RCT AM 74163-81-8

STAGE(1)

RGT D 6674-22-2 DBU SOL 75-05-8 MeCN

STAGE(2)

RGT AC 90825-38-0 Phosphorus(1+), ethyltris(N-ethylethanaminato)-, (T-4)-

SOL 75-09-2 CH2Cl2

STAGE(3)

SOL 75-09-2 CH2C12

STAGE (4)

RCT AN 84793-24-8 SOL 75-09-2 CH2C12

PRO AO **85441-61-8**

L23 ANSWER 2 OF 4 CASREACT COPYRIGHT 2003 ACS

ACCESSION NUMBER:

135:288704 CASREACT

TITLE:

Preparation of stable pharmaceutical

formulations containing moexipril magnesium and

the preparation of moexipril magnesium

INVENTOR(S):

Sherman, Bernard Charles

PATENT ASSIGNEE(S):

Can.

SOURCE:

Eur. Pat. Appl., 7 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

EP 1142878 A2 20011010 EP 2001-302450 20010316

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,

PT, IE, SI, LT, LV, FI, RO
US 2002037858 A1 20020328 US 2001-809173 20010316
PRIORITY APPLN. INFO.: CA 2000-2303481 20000405
AB Moexipril magnesium, a more stable form of moexipril, is prepd. by reacting moexipril or one of its acid-addn. salts (e.g., moexipril hydrochloride) with an alk. magnesium compd. (e.g., magnesium hydroxide) in the presence of a solvent (e.g., water and acetone) to convert most or all of the moexipril or the moexipril acid addn. salt into moexipril magnesium.

RX(1) OF 1 A ===> B

HCl

Α

$$\xrightarrow{(1)}$$

●1/2 Mg

Ŕ

RX(1) RCT A 82586-52-5 PRO B 365278-21-3 SOL 7732-18-5 Water, 67-64-1 Me2CO

L23 ANSWER 3 OF 4 CASREACT COPYRIGHT 2003 ACS

Searcher :

Shears

308-4994

ACCESSION NUMBER:

117:172104 CASREACT

TITLE:

Methods for the synthesis of aminosuberic acid

derivatives

INVENTOR(S):

Hoffmann, Gerhard; Liedtke, Bernhard; Vollmer,

Karl Otto

PATENT ASSIGNEE(S):

SOURCE:

Goedecke AG, Germany Ger. Offen., 6 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 4037960	A1	19920604	DE 1990-4037960	19901129
PRIORITY APPLN. INFO.	:		DE 1990-4037960	19901129

OTHER SOURCE(S):

MARPAT 117:172104

GI For diagram(s), see printed CA Issue.

Aminosuberic acid derivs. I (R1 = alkyl or alkenyl with up to 6carbon atoms, C5-7-cycloalkyl, C5-7-cycloalkenyl, C7-12-cycloalkylalkyl, C6-10-aryl, C7-14-aralkyl, mono- or bicyclic heterocyclic group; R2 = aryl; R3 = C1-6-alkyl, C2-6-alkenyl, C7-14-aralkyl; Z forms a heterocyclic ring) were prepd. by the selective cleavage of PhCH2O2CCHP1NHCH(CO2R3)CH2COR2 with AlCl3, condensing the resulting HO2CCHR1NHCH(CO2R3)CH2COR2 with heterocyclic compd. II, and hydrogenating the resulting ketone III with H2, deuterium or tritium. Thus, (S,S)-PhCOCH2CH(CO2Et)-Ala-OCH2Ph was debenzylated with AlCl3 in the presence of anisole in CH2Cl2/MeNO2 to give 86% (S,S)-PhCOCH2CH(CO2Et)-Ala-OH, which was condensed with (S)-1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid benzyl ester by DCC/1-hydroxybenzotriazole in CH2Cl2 to give product IV (isolated as the HCl salt). IV was hydrogenated over Pd/C in the presence of HCl to give 88% tetrahydroisoquinoline V.HCl.

RX(3) OF 6 L ===> M...

> 308-4994 Searcher : Shears

HCl

L

$$\stackrel{(3)}{\longrightarrow}$$

HC1

M YIELD 60%

RX(3) RCT L 143381-50-4

RGT N 1333-74-0 H2, O 7647-01-0 HCl

PRO M **82586-55-8**CAT 7440-05-3 Pd
SOL 60-29-7 Et20

L23 ANSWER 4 OF 4 CASREACT COPYRIGHT 2003 ACS

ACCESSION NUMBER:

105:208744 CASREACT

TITLE:

Synthesis of novel angiotensin converting enzyme inhibitor quinapril and related compounds. A divergence of structure-activity relationships

for non-sulfhydryl and sulfhydryl types AUTHOR(S):

Klutchko, Sylvester; Blankley, C. John; Fleming,

Robert W.; Hinkley, Jack M.; Werner, Ann E.; Nordin, Ivan; Holmes, Ann; Hoefle, Milton L.;

Cohen, David M.; et al.

Dep. Chem., Warner-Lambert/Parke-Davis Pharm. CORPORATE SOURCE:

Res., Ann Arbor, MI, 48106, USA

Journal of Medicinal Chemistry (1986), 29(10), SOURCE:

1953-61

CODEN: JMCMAR; ISSN: 0022-2623

I

DOCUMENT TYPE: Journal

LANGUAGE: English

GT

PhCH2CH2CH (
$$CO_2R^1$$
) NHCHMeCON RO2C

The synthesis and angiotensin-converting enzyme (ACE) inhibiting AΒ activities of quinapril (S,S,S)-I (R=R2=H,R1=Et), its active diacid (S,S,S)-I (R=R1=R2=H), and its dimethoxy analog (S,S,S)-I (R = H, R1 = Et, R2 = MeO) are reported. Thus, (S)-1,2,3,4-tetrahydro-3-isoquinolinecarboxylic acid 1,1-dimethylethyl ester was acylated with (S,S)-PhCH2CH2CH(CO2Et)NHCHMeCO2H followed by hydrolysis of the product to give (S,S,S)-I (R=R2=H,R1=Et). These tetrahydro-3isoquinolinecarboxylic acid derivs. possess in vitro potency and in vivo efficacy equiv. to that of enalapril. Sulfhydryl analogs with the same structural variation are also highly potent. In contrast, tetrahydro-1-isoquinolinecarboxylic acid and homologous isoindoline-1-carboxylic acid analogs show a striking divergence in potency between the two types, sulfhydryl analogs being essentially inactive, while non-sulfhydryl analogs are equipotent with the proline prototype. This is the first evidence suggesting that alternate binding modes may exist for the two major structural classes of small mol. ACE inhibitors.

RX(5) OF 91 ...T ===>

● HCl

Т

U: CM 2 ·

RX(5) RCT T 82586-55-8 RGT F 7647-01-0 HC1 PRO U 103733-36-4

SOL 64-17-5 EtOH

(FILE 'DJSMDS, CHEMINFORMRX, CHEMREACT' ENTERED AT 12:10:38 ON 21 APR 2003)

L21 STR

Searcher :

Shears

308-4994

REP G1=(1-3) CH2
VAR G2=OH/30
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS UNS AT 32
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RSPEC I NUMBER OF NODES IS 32

STEREO ATTRIBUTES: NONE L24 0 SEA L21

FILE 'HOME' ENTERED AT 12:11:21 ON 21 APR 2003